(FILE 'HOME' ENTERED AT 15:45:30 ON 29 SEP 2003)

FILE 'BIOSIS, CAPLUS, MEDLINE, WPIDS, EMBASE, SCISEARCH' ENTERED AT 15:45:38 ON 29 SEP 2003

E SCHUBERT

E E3 AND WALTER

E SCHUBERT

L1 1 S E3 AND WALTER

1 S E3 AND WALTER

229 S E3 AND CELL

L45 S L3 AND TARGET

FILE 'STNGUIDE' ENTERED AT 15:52:18 ON 29 SEP 2003

O S CELL SPECIFIC TARGET AND PROCESS

0 S CELL SPECIFIC TARGET L6

L70 S CELL- SPECIFIC TARGET

0 S CELL- SPECIFIC AND SCHUBERT rs

0 S MELK L9

L10 4 S PROTEOMICS

> FILE 'BIOSIS, CAPLUS, MEDLINE, WPIDS, EMBASE, SCISEARCH' ENTERED AT 16:06:49 ON 29 SEP 2003

FILE 'STNGUIDE' ENTERED AT 16:07:16 ON 29 SEP 2003

FILE 'BIOSIS, CAPLUS, MEDLINE, WPIDS, EMBASE, SCISEARCH' ENTERED AT 16:07:32 ON 29 SEP 2003

16796 S PROTEOMIC? L11

L12 1 S L11 AND MELK

L13 2438 S PROTEOMIC AND PROCESS

8 S L13 AND CULTIVATED

1282 S L13 AND CELL

L16 0 S L15 AND SCHUBERT

L17 15 S L15 AND HETEROGENEOUS

L18 1 S L15 AND PROTEIN CHIPS

L19 4 S L15 AND AUTOMATIC?

FILE 'STNGUIDE' ENTERED AT 16:21:29 ON 29 SEP 2003

FILE 'BIOSIS, CAPLUS, MEDLINE, WPIDS, EMBASE, SCISEARCH' ENTERED AT 16:25:49 ON 29 SEP 2003

L20 27 S L15 AND ENDOTHELI?

L21 16 DUP REM L20 (11 DUPLICATES REMOVED)

=>

L2

L3

L5

Search Report

89/808, 225

FILE 'HOME' ENTERED AT 15:45:30 ON 29 SEP 2003

=> FIL BIOSIS, CAPLUS, MEDLINE, WPIDS, EMBASE, SCISEARCH COST IN U.S. DOLLARS SINCE FILE

ENTRY

TOTAL SESSION

FULL ESTIMATED COST

0.21

0.21

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FILE 'SCISEARCH' ENTERED AT 15:45:38 ON 29 SEP 2003 COPYRIGHT 2003 THOMSON ISI

=> e E1 E2 E3 E4 E5 E6 E7 E8 E9 E10 E11 E12	schubert 69 1 1378> 1 33 2 4 2 2 2 22 33 1	SCHUBERGI/BI SCHUBERL/BI SCHUBERT/BI SCHUBERTA/BI SCHUBERTELLA/BI SCHUBERTELLID/BI SCHUBERTELLIDAE/BI SCHUBERTELLINAE/BI SCHUBERTELLINIDS/BI SCHUBERTH/BI SCHUBERTI/BI SCHUBERTIA/BI
=> e E1 E2 E3 E4 E5 E6 E7 E8 E9 E10 E11		E2ZZ/BI E3/BI E3 AND WALTER/BI E3.0/BI E3.10/BI E3.10.4K/BI E3.2.1.21/BI E3.3/BI E3.39.214/BI E3.4/BI E3.4/BI E3.4/BI E3.5/BI
=> e E1 E2 E3 E4 E5 E6	schubert 69 1 1378> 1 33 2	SCHUBERGI/BI SCHUBERL/BI SCHUBERT/BI SCHUBERTA/BI SCHUBERTELLA/BI SCHUBERTELLID/BI

4

SCHUBERTELLIDAE/BI

E7

E8 2 SCHUBERTELLINAE/BI E9 2 SCHUBERTELLINIDS/BI E10 22 SCHUBERTH/BI 33 E11 SCHUBERTI/BI E12 1 SCHUBERTIA/BI => s e3 and walter 1 SCHUBERT/BI AND WALTER => d l1 ibib abs ANSWER 1 OF 1 MEDLINE on STN ACCESSION NUMBER: 84269013 MEDLINE DOCUMENT NUMBER: 84269013 PubMed ID: 6748618 TITLE: Psychometric principles in the selection, interpretation, and evaluation of communication self-assessment inventories. Demorest M E; Walden B E AUTHOR: JOURNAL OF SPEECH AND HEARING DISORDERS, (1984 Aug) 49 (3) SOURCE: 226-40. Journal code: 0376335. ISSN: 0022-4677. PUB. COUNTRY: United States DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE) LANGUAGE: English Priority Journals FILE SEGMENT: ENTRY MONTH: 198409 ENTRY DATE: Entered STN: 19900320 Last Updated on STN: 19900320 Entered Medline: 19840905 AB A variety of self-assessment inventories have been introduced in recent years for use with hearing-impaired patients. These instruments differ considerably, both conceptually and operationally. Audiologists, therefore, are faced with the task of selecting a test instrument that is appropriate to their patient population and testing purpose. outlines the psychometric principles that guide the selection, interpretation, and evaluation of self-assessment inventories. application of these principles to a specific clinical population is illustrated by three studies of the Hearing Performance Inventory (Giolas, Owens, Lamb, & Schubert, 1979) conducted at Walter Reed Army Medical Center (WRAMC). => s e3 and walter L2 1 SCHUBERT/BI AND WALTER => s e3 and cell 229 SCHUBERT/BI AND CELL => s 13 and target 5 L3 AND TARGET => d 14 1-5 ibib abs ANSWER 1 OF 5 CAPLUS COPYRIGHT 2003 ACS on STN ACCESSION NUMBER: 1999:432073 CAPLUS DOCUMENT NUMBER: 131:75469 TITLE: Convection in the presence of a first-order phase change Sakurai, Shinichi; Tschammer, Armin; Pesch, Werner; AUTHOR(S): Ahlers, Guenter CORPORATE SOURCE: Department of Physics and Center for Nonlinear

Science, University of California at Santa Barbara,

Santa Barbara, CA, 93106, USA

SOURCE: Physical Review E: Statistical Physics, Plasmas,

Fluids, and Related Interdisciplinary Topics (1999),

60(1), 539-550

CODEN: PLEEE8; ISSN: 1063-651X

PUBLISHER: American Physical Society

DOCUMENT TYPE: Journal LANGUAGE: English

We report exptl. and theor. results for two-phase convection in a thin ΑB horizontal layer of a fluid with a first-order phase change and heated from below. A top layer of the nematic phase of a liq. crystal is located above the bottom layer of the isotropic phase of the same substance. A horizontal field of 1000 G is applied to align the director of the nematic phase. Over some ranges of the thickness of the isotropic phase, and in sufficiently large thermal gradients, the more dense nematic phase can be stably stratified above the less dense isotropic one, with a stable interface between them. Based on the equations of motion derived for this problem by Busse and Schubert [J. Fluid Mech. 46, 801 (1971)], we evaluate the bifurcation lines between the quiescent and convecting states and the corresponding crit. wave vectors as a function of the interface position. We report exptl. measurements based on Nusselt-no. detns. for the locations of the bifurcation lines. They are in good agreement with the theor. results. We also report approx. detns. of the crit. wave nos. which are semiquant. consistent with the theory. A great diversity of patterns is obsd. in the convecting states, including normal and parallel rolls, rolls with defects and disorder, target patterns and spirals, and cellular flow with upflow or downflow at the cell center. These patterns are discussed in terms of the breaking of the mirror symmetry at the horizontal midplane by the

interface, and in terms of the orienting effects of the magnetic field.

REFERENCE COUNT: 40 THERE ARE 40 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 2 OF 5 CAPLUS COPYRIGHT 2003 ACS on STN

ACCESSION NUMBER: 1977:565312 CAPLUS

DOCUMENT NUMBER: 87:165312

TITLE: Selective **cell** adhesion of neuronal

cell lines

AUTHOR(S): Santala, Roger; Gottlieb, David I.; Littman, Daniel;

Glaser, Luis

CORPORATE SOURCE: Dep. Biol. Chem., Washington Univ., St. Louis, MO, USA

SOURCE: Journal of Biological Chemistry (1977), 252(21),

7625-34

CODEN: JBCHA3; ISSN: 0021-9258

DOCUMENT TYPE: Journal LANGUAGE: English

Cloned neural cell lines derived from ethylnitrosourea-treated rat embryos (Schubert, D., et al, 1974) adhered preferentially to monolayers of cells obtained by dissocn. of neural tissue of either chick or rat embryos. One such cloned line, B103, will bind to cells obtained from any of the major regions of the embryonal nervous system, but bound only poorly to other types of cells. A plasma membrane-enriched fraction prepd. from B103 cells showed the same relative binding characteristics to embryonal neural and nonneural cells as intact B103 cells. Treatment of the membranes with trypsin at low concns. or treatment of the target cells with low concns. of glutaraldehyde or HCHO also abolished binding. A very similar binding pattern to that of B103 cells and plasma membranes was shown by B50 and B65 cells and plasma membranes in that both of these cell lines bound preferentially to monolayers prepd. from cells from embryonal nervous tissue. The plasma membranes from these cells however

showed significant differences in binding to other cultured neural cell lines. Apparently, only a part of the cells adhesive components is retained on the isolated plasma membrane, and there may be several adhesive components on each cell type. The cloned neural cells are apparently a suitable model system for the study of selective cell adhesion.

L4 ANSWER 3 OF 5 MEDLINE on STN

ACCESSION NUMBER: 2002244253 MEDLINE

DOCUMENT NUMBER: 21966828 PubMed ID: 11969793

TITLE: Convection in the presence of a first-order phase change.

AUTHOR: Sakurai S; Tschammer A; Pesch W; Ahlers G

CORPORATE SOURCE: Department of Physics and Center for Nonlinear Science,

University of California at Santa Barbara, Santa Barbara,

California 93106, USA.

SOURCE: PHYSICAL REVIEW. E, STATISTICAL PHYSICS, PLASMAS, FLUIDS,

AND RELATED INTERDISCIPLINARY TOPICS, (1999 Jul) 60 (1)

539-50.

Journal code: 9887340. ISSN: 1063-651X.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English ENTRY MONTH: 200208

ENTRY DATE: Entered STN: 20020502

Last Updated on STN: 20020829 Entered Medline: 20020827

We report experimental and theoretical results for two-phase convection in AΒ a thin horizontal layer of a fluid with a first-order phase change and heated from below. A top layer of the nematic phase of a liquid crystal is located above the bottom layer of the isotropic phase of the same substance. A horizontal field of 1000 G is applied in order to align the director of the nematic phase. Over some ranges of the thickness of the isotropic phase, and in sufficiently large thermal gradients, the more dense nematic phase can be stably stratified above the less dense isotropic one, with a stable interface between them. Based on the equations of motion derived for this problem by Busse and Schubert [J. Fluid Mech. 46, 801 (1971)], we evaluate the bifurcation lines between the quiescent and convecting states and the corresponding critical wave vectors as a function of the interface position. We report experimental measurements based on Nusselt-number determinations for the locations of the bifurcation lines. They are in good agreement with the theoretical results. We also report approximate determinations of the critical wave numbers which are semiquantitatively consistent with the theory. A great diversity of patterns is observed in the convecting states, including normal and parallel rolls, rolls with defects and disorder, target patterns and spirals, and cellular flow with upflow or downflow at the cell center. These patterns are discussed in terms of the breaking of the mirror symmetry at the horizontal midplane by the interface, and in terms of the orienting effects of the magnetic field.

L4 ANSWER 4 OF 5 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED.

on STN

ACCESSION NUMBER: 78229198 EMBASE

DOCUMENT NUMBER: 1978229198

TITLE: Selective cell adhesion of neuronal cell

lines.

AUTHOR: Santala R.; Gottlieb D.I.; Littman D.; Glaser L.

CORPORATE SOURCE: Div. Biol. Biomed. Sci., Dept. Biol. Chem., Washington

Univ., St Louis, Mo. 63110, United States

SOURCE: Journal of Biological Chemistry, (1977) 252/21 (7625-7634).

CODEN: JBCHA3

COUNTRY: United States

DOCUMENT TYPE: Journal

FILE SEGMENT: Neurology and Neurosurgery 800

> 029 Clinical Biochemistry

LANGUAGE: English

Cloned neural cell lines derived from ethylnitrosourea-treated rat embryos (Schubert, D., Heinemann, S., Carlisle, W., Tarikas,

H., Kimes, B., Patrick, J., Steinbach, J. H., Culp, W., and Brandt, B. L. (1974) Nature 249, 224-227) adhere preferentially to monolayers of cells obtained by dissociation of neural tissue of either chick or rat embryos. One such cloned line, B103, has been investigated in some

detail. B103 cells will bind to cells obtained from

any of the major regions of the embryonal nervous system. B103 cells will bind only poorly to chick embryo fibroblasts, Chinese

hamster ovary cells, or embryonal liver cells from

either the chick or the rat. A plasma membrane-enriched fraction prepared from B103 cells shows the same relative binding characteristics

to embryonal neural and non-neural cells as intact B103 cells. Treatment of the membranes with trypsin at low concentrations or treatment of the target cells with

low concentrations of glutaraldehyde or formaldehyde also abolishes binding. Binding does not take place at 0.degree.. A very similar binding pattern to that of B103 cells and plasma membranes is shown by B50 and B65 cells and plasma membranes in that both of these

cell lines bind preferentially to monolayers prepared from cells from embryonal nervous tissue. The plasma membranes from these cells however show significant differences in binding to other cultured neural cell lines. It is suggested that only a part of the cells adhesive components is retained on the

isolated plasma membrane and that there may be several adhesive components on each cell type. The cloned neural cells appear to

be a suitable model system for the study of selective cell adhesion.

ANSWER 5 OF 5 SCISEARCH COPYRIGHT 2003 THOMSON ISI on STN

ACCESSION NUMBER: 1999:589964 SCISEARCH

THE GENUINE ARTICLE: 218KC

TITLE: Convection in the presence of a first-order phase change

AUTHOR: Sakurai S (Reprint); Tschammer A; Pesch W; Ahlers G CORPORATE SOURCE: UNIV CALIF SANTA BARBARA, DEPT PHYS, SANTA BARBARA, CA

93106 (Reprint); UNIV CALIF SANTA BARBARA, CTR NONLINEAR SCI, SANTA BARBARA, CA 93106; KYOTO INST TECHNOL, DEPT POLYMER SCI & ENGN, SAKYO KU, KYOTO 6068585, JAPAN; UNIV

BAYREUTH, INST PHYS, D-95440 BAYREUTH, GERMANY

COUNTRY OF AUTHOR: USA; JAPAN; GERMANY

PHYSICAL REVIEW E, (JUL 1999) Vol. 60, No. 1, pp. 539-550. SOURCE:

Publisher: AMERICAN PHYSICAL SOC, ONE PHYSICS ELLIPSE,

COLLEGE PK, MD 20740-3844.

ISSN: 1063-651X.

DOCUMENT TYPE: Article; Journal

FILE SEGMENT: PHYS LANGUAGE: English REFERENCE COUNT: 37

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

AB We report experimental and theoretical results for two-phase convection in a thin horizontal layer of a fluid with a first-order phase change and heated from below. A top layer of the nematic phase of a liquid crystal is located above the bottom layer of the isotropic phase of the same. substance. A horizontal field of 1000 G is applied in order to align the director of the nematic phase. Over some ranges of the thickness of the isotropic phase, and in sufficiently large thermal gradients, the more dense nematic phase can be stably stratified above the less dense

isotropic one, with a stable interface between them. Based on the equations of motion derived for this problem by Busse and Schubert [J. Fluid Mech. 46, 801 (1971)], we evaluate the bifurcation lines between the quiescent and convecting states and the corresponding critical wave vectors as a function of the interface position. We report experimental measurements based on Nusselt-number determinations for the locations of the bifurcation lines. They are in good agreement with the theoretical results. We also report approximate determinations of the critical wave numbers which are semiquantitatively consistent with the theory. A great diversity of patterns is observed in the convecting states, including normal and parallel rolls, rolls with defects and disorder, target patterns and spirals, and cellular flow with upflow or downflow at the cell center. These patterns are discussed in terms of the breaking of the mirror symmetry at the horizontal midplane by the interface, and in terms of the orienting effects of the magnetic field. [S1063-651X(99)05707-41.

=> FIL STNGUIDE COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION 35.20 FULL ESTIMATED COST 34.99 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION -1.30CA SUBSCRIBER PRICE -1.30

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FILE CONTAINS CURRENT INFORMATION.
LAST RELOADED: Sep 26, 2003 (20030926/UP).

=> s cell specific target and process

16 CELL

29 SPECIFIC

20 TARGET

O CELL SPECIFIC TARGET

(CELL(W)SPECIFIC(W)TARGET)

14 PROCESS

14 PROCESSES

23 PROCESS

(PROCESS OR PROCESSES)

L5 0 CELL SPECIFIC TARGET AND PROCESS

=> s cell specific target

16 CELL

29 SPECIFIC

20 TARGET

L6 0 CELL SPECIFIC TARGET

(CELL(W) SPECIFIC(W) TARGET)

=> s cell- specific target

16 CELL

29 SPECIFIC

20 TARGET

L7 0 CELL- SPECIFIC TARGET

(CELL(W)SPECIFIC(W)TARGET)

=> s cell- specific and schubert

16 CELL 29 SPECIFIC 0 CELL- SPECIFIC (CELL(W)SPECIFIC) 0 SCHUBERT L8 0 CELL- SPECIFIC AND SCHUBERT => s MELK 0 MELK L9 0 MELK => s proteomics L10 4 PROTEOMICS => d llo 1-4 ibib abs 'L10' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE' 'IBIB' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE' 'ABS' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE' The following are valid formats: The default display format is GEN. GEN --- AN, DBN, DESC, PROD DBS --- AN, DBN, DESC, SUBJ, SI, DATA, UA, PROD, SUPP, REPR, SFIELD, PROP, DFIELD, TFIELD, FEAT, EXAMPLE ALL --- AN, DBN, DESC, SUBJ, CC, SI, DATA, UA, NTE, PROD, SUPP, REPR, SFIELD, PROP, DFIELD, TFIELD, FEAT, PRICE, EXAMPLE COVERAGE - AN, DBN, SUBJ, CC, SI, DATA FIELDS --- AN, DBN, SFIELD, PROP, DFIELD, TFIELD HIT ----- All fields containing hit terms KWIC ---- All fields containing hit terms OCC ----- List of display fields containing hit terms Hit terms will be highlighted in all displayable fields. To display a particular field or fields, enter the display field codes. For a list of display field codes, enter 'HELP DFIELDS' at an arrow prompt (=>). Examples of formats include: 'GEN'; 'AN'; 'SI,CC'. You may specify the format fields in any order, and the information will be displayed in the same order as the format specification. The same formats (except for HIT, KWIC, and OCC) may be used with the DISPLAY ACC command to display the record for a specified Accession Number. ENTER DISPLAY FORMAT (GEN):s proteomicsend 'S' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE' 'PROTEOMICSEND' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE' The following are valid formats: The default display format is GEN. GEN --- AN, DBN, DESC, PROD DBS --- AN, DBN, DESC, SUBJ, SI, DATA, UA, PROD, SUPP, REPR, SFIELD, PROP, DFIELD, TFIELD, FEAT, EXAMPLE ALL --- AN, DBN, DESC, SUBJ, CC, SI, DATA, UA, NTE,

PROD, SUPP, REPR, SFIELD, PROP, DFIELD, TFIELD, FEAT, PRICE, EXAMPLE COVERAGE - AN, DBN, SUBJ, CC, SI, DATA

FIELDS --- AN, DBN, SFIELD, PROP, DFIELD, TFIELD HIT ----- All fields containing hit terms

KWIC ---- All fields containing hit terms

OCC ----- List of display fields containing hit terms

Hit terms will be highlighted in all displayable fields.

To display a particular field or fields, enter the display field codes. For a list of display field codes, enter 'HELP DFIELDS' at an arrow prompt (=>). Examples of formats include: 'GEN'; 'AN'; 'SI,CC'. You may specify the format fields in any order, and the information will be displayed in the same order as the format specification.

The same formats (except for HIT, KWIC, and OCC) may be used with the DISPLAY ACC command to display the record for a specified Accession Number.

ENTER DISPLAY FORMAT (GEN): d 110 1-4 ibib abs 'D' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE' 'L34' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE' '1-4' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE' 'IBIB' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE' 'ABS' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE'

The following are valid formats:

The default display format is GEN.

GEN --- AN, DBN, DESC, PROD

DBS --- AN, DBN, DESC, SUBJ, SI, DATA, UA, PROD, SUPP, REPR, SFIELD, PROP, DFIELD, TFIELD, FEAT, EXAMPLE

ALL --- AN, DBN, DESC, SUBJ, CC, SI, DATA, UA, NTE, PROD, SUPP, REPR, SFIELD, PROP, DFIELD, TFIELD, FEAT, PRICE, EXAMPLE

COVERAGE - AN, DBN, SUBJ, CC, SI, DATA FIELDS --- AN, DBN, SFIELD, PROP, DFIELD, TFIELD

HIT ---- All fields containing hit terms

KWIC ---- All fields containing hit terms

OCC ----- List of display fields containing hit terms

Hit terms will be highlighted in all displayable fields.

To display a particular field or fields, enter the display field codes. For a list of display field codes, enter 'HELP DFIELDS' at an arrow prompt (=>). Examples of formats include: 'GEN'; 'AN'; 'SI,CC'. You may specify the format fields in any order, and the information will be displayed in the same order as the format specification.

The same formats (except for HIT, KWIC, and OCC) may be used with the DISPLAY ACC command to display the record for a specified Accession Number.

ENTER DISPLAY FORMAT (GEN):d 110 ibib abs 'D' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE' 'L34' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE' 'IBIB' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE' 'ABS' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE'

The following are valid formats:

The default display format is GEN.

GEN --- AN, DBN, DESC, PROD

DBS --- AN, DBN, DESC, SUBJ, SI, DATA, UA, PROD, SUPP, REPR, SFIELD, PROP, DFIELD, TFIELD, FEAT, EXAMPLE

ALL --- AN, DBN, DESC, SUBJ, CC, SI, DATA, UA, NTE, PROD, SUPP, REPR, SFIELD, PROP, DFIELD, TFIELD, FEAT, PRICE, EXAMPLE

COVERAGE - AN, DBN, SUBJ, CC, SI, DATA

FIELDS --- AN, DBN, SFIELD, PROP, DFIELD, TFIELD

HIT ----- All fields containing hit terms

KWIC ---- All fields containing hit terms

OCC ----- List of display fields containing hit terms

Hit terms will be highlighted in all displayable fields.

To display a particular field or fields, enter the display field codes. For a list of display field codes, enter 'HELP DFIELDS' at an arrow prompt (=>). Examples of formats include: 'GEN'; 'AN'; 'SI,CC'. You may specify the format fields in any order, and the information will be displayed in the same order as the format specification.

The same formats (except for HIT, KWIC, and OCC) may be used with the DISPLAY ACC command to display the record for a specified Accession Number.

ENTER DISPLAY FORMAT (GEN):d hist
'D' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE'
'HIST' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE'

The following are valid formats:

The default display format is GEN.

GEN --- AN, DBN, DESC, PROD

DBS --- AN, DBN, DESC, SUBJ, SI, DATA, UA, PROD, SUPP, REPR, SFIELD, PROP, DFIELD, TFIELD, FEAT, EXAMPLE

ALL --- AN, DBN, DESC, SUBJ, CC, SI, DATA, UA, NTE, PROD, SUPP, REPR, SFIELD, PROP, DFIELD, TFIELD, FEAT, PRICE, EXAMPLE

COVERAGE - AN, DBN, SUBJ, CC, SI, DATA

FIELDS --- AN, DBN, SFIELD, PROP, DFIELD, TFIELD

HIT ----- All fields containing hit terms

KWIC ---- All fields containing hit terms

OCC ---- List of display fields containing hit terms

Hit terms will be highlighted in all displayable fields.

To display a particular field or fields, enter the display field codes. For a list of display field codes, enter 'HELP DFIELDS' at an arrow prompt (=>). Examples of formats include: 'GEN'; 'AN'; 'SI,CC'. You may specify the format fields in any order, and the information will be displayed in the same order as the format specification.

The same formats (except for HIT, KWIC, and OCC) may be used with the DISPLAY ACC command to display the record for a specified Accession Number.

ENTER DISPLAY FORMAT (GEN):end

=> d his

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FILE 'BIOSIS, CAPLUS, MEDLINE, WPIDS, EMBASE, SCISEARCH' ENTERED AT 15:45:38 ON 29 SEP 2003

E SCHUBERT

E E3 AND WALTER

E SCHUBERT

L1 1 S E3 AND WALTER

L2 1 S E3 AND WALTER

L3 229 S E3 AND CELL

L4 5 S L3 AND TARGET

FILE 'STNGUIDE' ENTERED AT 15:52:18 ON 29 SEP 2003

L5 0 S CELL SPECIFIC TARGET AND PROCESS

L6 0 S CELL SPECIFIC TARGET

L7 0 S CELL- SPECIFIC TARGET

L8 0 S CELL- SPECIFIC AND SCHUBERT

L9 0 S MELK

L10 4 S PROTEOMICS

=> d 110 ibib abs 1-4

'IBIB' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE'

'ABS' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE'

The following are valid formats:

The default display format is GEN.

GEN --- AN, DBN, DESC, PROD

DBS --- AN, DBN, DESC, SUBJ, SI, DATA, UA, PROD, SUPP, REPR, SFIELD, PROP, DFIELD, TFIELD, FEAT, EXAMPLE

ALL --- AN, DBN, DESC, SUBJ, CC, SI, DATA, UA, NTE, PROD, SUPP, REPR, SFIELD, PROP, DFIELD, TFIELD, FEAT, PRICE, EXAMPLE

COVERAGE - AN, DBN, SUBJ, CC, SI, DATA

FIELDS --- AN, DBN, SFIELD, PROP, DFIELD, TFIELD

HIT ----- All fields containing hit terms

KWIC ---- All fields containing hit terms

OCC ---- List of display fields containing hit terms

Hit terms will be highlighted in all displayable fields.

To display a particular field or fields, enter the display field codes. For a list of display field codes, enter 'HELP DFIELDS' at an arrow prompt (=>). Examples of formats include: 'GEN'; 'AN'; 'SI,CC'. You may specify the format fields in any order, and the information will be displayed in the same order as the format specification.

The same formats (except for HIT, KWIC, and OCC) may be used with the DISPLAY ACC command to display the record for a specified Accession Number.

ENTER DISPLAY FORMAT (GEN):end

=> d hist

L5

(FILE 'HOME' ENTERED AT 15:45:30 ON 29 SEP 2003)

FILE 'BIOSIS, CAPLUS, MEDLINE, WPIDS, EMBASE, SCISEARCH' ENTERED AT 15:45:38 ON 29 SEP 2003

E SCHUBERT

E E3 AND WALTER

E SCHUBERT

L1 1 S E3 AND WALTER

L2 1 S E3 AND WALTER

L3 229 S E3 AND CELL

L4 5 S L3 AND TARGET

FILE 'STNGUIDE' ENTERED AT 15:52:18 ON 29 SEP 2003

0 S CELL SPECIFIC TARGET AND PROCESS

L6 0 S CELL SPECIFIC TARGET

L7 0 S CELL- SPECIFIC TARGET

L8 0 S CELL- SPECIFIC AND SCHUBERT

L9 0.S MELK

L10 4 S PROTEOMICS

=> FIL BIOSIS, CAPLUS, MEDLINE, WPIDS, EMBASE, SCISEARCH

COST IN U.S. DOLLARS SINCE FILE TOTAL

ENTRY SESSION

FULL ESTIMATED COST 1.44 36.64

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION

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=> d 110 abs ibib 1-4

YOU HAVE REQUESTED DATA FROM FILE 'STNGUIDE' - CONTINUE? (Y)/N:y

'ABS' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE'

'IBIB' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE'

The following are valid formats:

The default display format is GEN.

GEN --- AN, DBN, DESC, PROD
DBS --- AN, DBN, DESC, SUBJ, SI, DATA, UA, PROD,
SUPP, REPR, SFIELD, PROP, DFIELD, TFIELD,
FEAT, EXAMPLE
ALL --- AN, DBN, DESC, SUBJ, CC, SI, DATA, UA, NTE,
PROD, SUPP, REPR, SFIELD, PROP, DFIELD, TFIELD,
FEAT, PRICE, EXAMPLE
COVERAGE - AN, DBN, SUBJ, CC, SI, DATA
FIELDS --- AN, DBN, SFIELD, PROP, DFIELD, TFIELD
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KWIC ---- All fields containing hit terms
OCC ----- List of display fields containing hit terms

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ENTER DISPLAY FORMAT (GEN): FIL BIOSIS, CAPLUS, MEDLINE, WPIDS, EMBASE, SCISEARCH 'FIL' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE' 'BIOSIS' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE' 'CAPLUS' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE' 'MEDLINE' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE' 'WPIDS' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE' 'EMBASE' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE' 'SCISEARCH' IS NOT A VALID FORMAT FOR FILE 'STNGUIDE'

The following are valid formats:

GEN --- AN, DBN, DESC, PROD

The default display format is GEN.

DBS --- AN, DBN, DESC, SUBJ, SI, DATA, UA, PROD,
SUPP, REPR, SFIELD, PROP, DFIELD, TFIELD,
FEAT, EXAMPLE

ALL --- AN, DBN, DESC, SUBJ, CC, SI, DATA, UA, NTE,
PROD, SUPP, REPR, SFIELD, PROP, DFIELD, TFIELD,
FEAT, PRICE, EXAMPLE

COVERAGE - AN, DBN, SUBJ, CC, SI, DATA
FIELDS --- AN, DBN, SFIELD, PROP, DFIELD, TFIELD
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KWIC ---- All fields containing hit terms

OCC ----- List of display fields containing hit terms

Hit terms will be highlighted in all displayable fields.

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The same formats (except for HIT, KWIC, and OCC) may be used with the DISPLAY ACC command to display the record for a specified Accession Number.

ENTER DISPLAY FORMAT (GEN):end

=> s proteomic?

L11 16796 PROTEOMIC?

=> s 111 and MELK

L12 1 L11 AND MELK

=> d 112 abs ibib

L12 ANSWER 1 OF 1 MEDLINE on STN

MELK is an ultrasensitive topological proteomics AB technology analysing proteins on the single cell level (Multi-Epitope-Ligand-'Kartographie'). It can trace out large scale protein patterns with subcellular resolution, mapping the topological position of many proteins simultaneously in a cell. Thereby, it addresses higher level order in a proteome, referred to as the toponome, coding cell functions by topologically and timely determined webs of interacting proteins. The resulting cellular protein maps provide new structures in the proteome: single combinatorial protein patterns (s-CPP), and combinatorial protein pattern motifs (CPP-motifs), bound to superior units. They are images of functional protein networks, which are specific signatures of tissues, cell types, cell states and diseases. The technology unravels hierarchies of proteins related to particular cell functions or dysfunctions, thus identifying and prioritising key proteins within cell and tissue protein networks. Interlocking MELK with the drug screening machinery provides new clues related to the selection of target proteins, and functionally relevant hits and drug leads. present chapter summarizes the steps that have contributed to the establishment of the technology.

ACCESSION NUMBER: 2003397091 MEDLINE

DOCUMENT NUMBER: 22815613 PubMed ID: 12934931

TITLE: Topological proteomics, toponomics, MELK

-technology.

AUTHOR: Schubert Walter

CORPORATE SOURCE: MelTec Ltd., ZENIT-Building, Leipziger Strasse 44, 39120

Magdeburg, Germany.. info@meltec.de

SOURCE: ADVANCES IN BIOCHEMICAL ENGINEERING/BIOTECHNOLOGY, (2003)

83 189-209. Ref: 30

Journal code: 8307733. ISSN: 0724-6145. Germany: Germany, Federal Republic of Journal; Article; (JOURNAL ARTICLE)

General Review; (REVIEW)

(REVIEW, TUTORIAL)

LANGUAGE: English

PUB. COUNTRY:

DOCUMENT TYPE:

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200309

ENTRY DATE: Entered STN: 20030826

Last Updated on STN: 20030928 Entered Medline: 20030926

=> s proteomic and process

L13 2438 PROTEOMIC AND PROCESS

=> s 113 and cultivated

L14 8 L13 AND CULTIVATED

=> d l14 ibib abs

L14 ANSWER 1 OF 8 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN

ACCESSION NUMBER: 2003:232833 BIOSIS DOCUMENT NUMBER: PREV200300232833

TITLE: Proteomic analysis of Lactococcus lactis, a

lactic acid bacterium.

AUTHOR(S): Guillot, Alain; Gitton, Christophe; Anglade, Patricia;

Mistou, Michel-Yves (1)

CORPORATE SOURCE: (1) Unite de Biochimie et Structure des Proteines, Institut

National de la Recherche Agronomique, 78352, Jouy-en-Josas

cedex, France: mistou@jouy.inra.fr France

SOURCE: Proteomics, (March 2003, 2003) Vol. 3, No. 3, pp. 337-354.

print.

ISSN: 1615-9853.

DOCUMENT TYPE: Article LANGUAGE: English

Lactococcus lactis is a Gram-positive bacteria, which belongs to the group AΒ of lactic acid bacteria among which several genera play an essential role in the manufacture of food products. Cytosolic proteins of L. lactis IL1403 cultivated in M17 broth have been resolved by two-dimensional gel electrophoresis using two pH gradients (pH 4-7, 4.5-5.5). More than 230 spots were identified by peptide mass fingerprints, corresponding to 25% of the predicted acid proteome. The present study made it possible to describe at the proteome level a significant number of cellular pathways (glycolysis, fermentation, nucleotide metabolism, proteolysis, fatty acid and peptidoglycan synthesis) related to important physiological processes and technological properties. It also indicated that the fermentative metabolism, which characterizes L. lactis is associated with a high expression of glycolytic enzymes. Thirty-four proteins were matched to open reading frames for which there is no assigned function. The comparison at the proteome level of two strains of L. lactis showed an important protein polymorphism. The comparison of the proteomes of glucose- and lactose-grown cells revealed an unexpected link between the nature of the carbon source and the metabolism of pyrimidine nucleotides.

=> d 114 ibib abs 2-8

L14 ANSWER 2 OF 8 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN

ACCESSION NUMBER: 2001:292643 BIOSIS DOCUMENT NUMBER: PREV200100292643

TITLE: Genomic and proteomic analysis of microbial

function in the gastrointestinal tract of ruminants.

AUTHOR(S): White, Bryan A. (1); Morrison, Mark

CORPORATE SOURCE: (1) Departments of Animal Sciences and Veterinary

Pathobiology, and the Division of Nutritional Sciences, University of Illinois at Urbana-Champaign, 1207 West Gregory Drive, Urbana, IL, 61801: b-white2@uiuc.edu,

morrison.234@osu.edu USA

SOURCE: Asian-Australasian Journal of Animal Sciences, (June, 2001)

Vol. 14, No. 6, pp. 880-884. print.

ISSN: 1011-2367.

DOCUMENT TYPE: General Review

LANGUAGE: English SUMMARY LANGUAGE: English

AB Rumen microbiology research has undergone several evolutionary steps: the isolation and nutritional characterization of readily cultivated microbes; followed by the cloning and sequence analysis of individual genes relevant to key digestive processes; through to the use of small subunit ribosomal RNA (SSU rRNA) sequences for a

cultivation-independent examination of microbial diversity. Our knowledge of rumen microbiology has expanded as a result, but the translation of this information into productive alterations of ruminal function has been rather limited. For instance, the cloning and characterization of cellulase genes in Escherichia coli has yielded some valuable information about this complex enzyme system in ruminal bacteria. SSU rRNA analyses have also confirmed that a considerable amount of the microbial diversity in the rumen is not represented in existing culture collections. However, we still have little idea of whether the key, and potentially rate-limiting, gene products and (or) microbial interactions have been identified. Technologies allowing high throughput nucleotide and protein sequence analysis have led to the emergence of two new fields of investigation, genomics and proteomics. Both disciplines can be further subdivided into functional and comparative lines of investigation. The massive accumulation of microbial DNA and protein sequence data, including complete genome sequences, is revolutionizing the way we examine microbial physiology and diversity. We describe here some examples of our use of genomics- and proteomics-based methods, to analyze the cellulase system of Ruminococcus flavefaciens FD-1 and explore the genome of Ruminococcus albus 8. At Illinois, we are using bacterial artificial chromosome (BAC) vectors to create libraries containing large (>75 kbases), contiguous segments of DNA from R. flavefaciens FD-1. Considering that every bacterium is not a candidate for whole genome sequencing, BAC libraries offer an attractive, alternative method to perform physical and functional analyses of a bacterium's genome. Our first plan is to use these BAC clones to determine whether or not cellulases and accessory genes in R. flavefaciens exist in clusters of orthologous genes (COGs). Proteomics is also being used to complement the BAC library/DNA sequencing approach. Proteins differentially expressed in response to carbon source are being identified by 2-D SDS-PAGE, followed by in-gel-digests and peptide mass mapping by MALDI-TOF Mass Spectrometry, as well as peptide sequencing by Edman degradation. At Ohio State, we have used a combination of functional proteomics, mutational analysis and differential display RT-PCR to obtain evidence suggesting that in addition to a cellulosome-like mechanism, R. albus 8 possesses other mechanisms for adhesion to plant surfaces. Genome walking on either side of these differentially expressed transcripts has also resulted in two interesting observations: i) a relatively large number of genes with no matches in the current databases and; ii) the identification of genes with a high level of sequence identity to those identified, until now, in the archaebacteria. Genomics and proteomics will also accelerate our understanding of microbial interactions, and allow a greater degree of in situ analyses in the future. The challenge is to utilize genomics and proteomics to improve our fundamental understanding of microbial physiology, diversity and ecology, and overcome constraints to ruminal function.

L14 ANSWER 3 OF 8 CAPLUS COPYRIGHT 2003 ACS on STN

ACCESSION NUMBER: 2003:244402 CAPLUS

DOCUMENT NUMBER: 139:130342

TITLE: Proteomic analysis of Lactococcus lactis, a

lactic acid bacterium

AUTHOR(S): Guillot, Alain; Gitton, Christophe; Anglade, Patricia;

Mistou, Michel-Yves

CORPORATE SOURCE: Institut National de la Recherche Agronomique,

Jouy-en-Josas, Fr.

SOURCE: Proteomics (2003), 3(3), 337-354

CODEN: PROTC7; ISSN: 1615-9853 Wiley-VCH Verlag GmbH & Co. KGaA

DOCUMENT TYPE: Journal LANGUAGE: English

PUBLISHER:

AB Lactococcus lactis is a Gram-pos. bacteria, which belongs to the group of

lactic acid bacteria among which several genera play an essential role in the manuf. of food products. Cytosolic proteins of L. lactis IL1403 cultivated in M17 broth have been resolved by two-dimensional gel electrophoresis using two pH gradients (pH 4-7, 4.5-5.5). More than 230 spots were identified by peptide mass fingerprints, corresponding to 25% of the predicted acid proteome. The present study made it possible to describe at the proteome level a significant no. of cellular pathways (glycolysis, fermn., nucleotide metab., proteolysis, fatty acid and peptidoglycan synthesis) related to important physiol. processes and technol. properties. It also indicated that the fermentative metab., which characterizes L. lactis is assocd. With a high expression of glycolytic enzymes. Thirty-four proteins were matched to open reading frames for which there is no assigned function. The comparison at the proteome level of two strains of L. lactis showed an important protein polymorphism. The comparison of the proteomes of glucose- and lactose-grown cells revealed an unexpected link between the nature of the carbon source and the metab. of pyrimidine nucleotides.

REFERENCE COUNT: 54 THERE ARE 54 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L14 ANSWER 4 OF 8 CAPLUS COPYRIGHT 2003 ACS on STN

ACCESSION NUMBER: 2001:398436 CAPLUS

DOCUMENT NUMBER: 136:129446

TITLE: Genomic and **proteomic** analysis of microbial

function in the gastrointestinal tract of ruminants

AUTHOR(S): White, Bryan A.; Morrison, Mark

CORPORATE SOURCE: Departments of Animal Sciences and Veterinary

Pathobiology, University of Illinois at Urbana-Champaign, Urbana, IL, 61801, USA

SOURCE: Asian-Australasian Journal of Animal Sciences (2001),

14(6), 880-884

CODEN: AJASEL; ISSN: 1011-2367

PUBLISHER: Asian-Australasian Journal of Animal Sciences

DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

AΒ A review. Rumen microbiol. research has undergone several evolutionary steps: the isolation and nutritional characterization of readily cultivated microbes; followed by the cloning and sequence anal. of individual genes relevant to key digestive processes; through to the use of small subunit rRNA (SSU rRNA) sequences for a cultivation-independent examn. of microbial diversity. Our knowledge of rumen microbiol. has expanded as a result, but the translation of this information into productive alterations of ruminal function has been rather limited. For instance, the cloning and characterization of cellulase genes in Escherichia coli has yielded some valuable information about this complex enzyme system in ruminal bacteria. SSU rRNA analyses have also confirmed that a considerable amt. of the microbial diversity in the rumen is not represented in existing culture collections. However, we still have little idea of whether the key, and potentially rate-limiting, gene products and (or) microbial interactions have been identified. Technologies allowing high throughput nucleotide and protein sequence anal. have led to the emergence of two new fields of investigation, genomics and proteomics. Both disciplines can be further subdivided into functional and comparative lines of investigation. The massive accumulation of microbial DNA and protein sequence data, including complete genome sequences, is revolutionizing the way we examine microbial physiol. and diversity. We describe here some examples of our use of genomics- and proteomics-based methods, to analyze the cellulase system of Ruminococcus flavefaciens FD-1 and explore the genome of R. albus 8. At Illinois, we are using bacterial artificial chromosome (BAC) vectors to create libraries contg. large (>75 kbases), contiguous segments of DNA from R. flavefaciens FD-1. Considering that every bacterium is not

a candidate for whole genome sequencing, BAC libraries offer an attractive, alternative method to perform phys. and functional analyses of a bacterium's genome. Our first plan is to use these BAC clones to det. whether or not cellulases and accessory genes in R. flavefaciens exist in clusters of orthologous genes (COGs). Proteomics is also being used to complement the BAC library/DNA sequencing approach. Proteins differentially expressed in response to carbon source are being identified by 2-D SDS-PAGE, followed by in-gel-digests and peptide mass mapping by MALDI-TOF Mass Spectrometry, as well as peptide sequencing by Edman degrdn. At Ohio State, we have used a combination of functional proteomics, mutational anal. and differential display RT-PCR to obtain evidence suggesting that in addn. to a cellulosome-like mechanism, R. albus 8 possesses other mechanisms for adhesion to plant surfaces. Genome walking on either side of these differentially expressed transcripts has also resulted in two interesting observations: (i) a relatively large no. of genes with no matches in the current databases and; (ii) the identification of genes with a high level of sequence identity to those identified, until now, in the archaebacteria. Genomics and proteomics will also accelerate our understanding of microbial interactions, and allow a greater degree of in situ analyses in the future. The challenge is to utilize genomics and proteomics to improve our fundamental understanding of microbial physiol., diversity and ecol., and overcome constraints to ruminal function.

REFERENCE COUNT:

29 THERE ARE 29 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L14 ANSWER 5 OF 8 MEDLINE on STN

ACCESSION NUMBER: 2003114020 IN-PROCESS

DOCUMENT NUMBER: 22514603 PubMed ID: 12627387

TITLE: Proteomic analysis of Lactococcus lactis, a

lactic acid bacterium.

AUTHOR: Guillot Alain; Gitton Christophe; Anglade Patricia; Mistou

Michel-Yves

CORPORATE SOURCE: Institut National de la Recherche Agronomique,

Jouy-en-Josas, France.

SOURCE: Proteomics, (2003 Mar) 3 (3) 337-54.

Journal code: 101092707. ISSN: 1615-9853.

PUB. COUNTRY: Germany: Germany, Federal Republic of DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: IN-PROCESS; NONINDEXED; Priority Journals

ENTRY DATE: Entered STN: 20030311

Last Updated on STN: 20030311

AΒ Lactococcus lactis is a Gram-positive bacteria, which belongs to the group of lactic acid bacteria among which several genera play an essential role in the manufacture of food products. Cytosolic proteins of L. lactis IL1403 cultivated in M17 broth have been resolved by two-dimensional gel electrophoresis using two pH gradients (pH 4-7, 4.5-5.5). More than 230 spots were identified by peptide mass fingerprints, corresponding to 25% of the predicted acid proteome. present study made it possible to describe at the proteome level a significant number of cellular pathways (glycolysis, fermentation, nucleotide metabolism, proteolysis, fatty acid and peptidoglycan synthesis) related to important physiological processes and technological properties. It also indicated that the fermentative metabolism, which characterizes L. lactis is associated with a high expression of glycolytic enzymes. Thirty-four proteins were matched to open reading frames for which there is no assigned function. The comparison at the proteome level of two strains of L. lactis showed an important protein polymorphism. The comparison of the proteomes of glucose- and lactose-grown cells revealed an unexpected link between the nature of the carbon source and the metabolism of pyrimidine nucleotides.

L14 ANSWER 6 OF 8 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED.

on STN

ACCESSION NUMBER: 2003124019 EMBASE

Proteomic analysis of Lactococcus lactis, a TITLE:

lactic acid bacterium.

AUTHOR: Guillot A.; Gitton C.; Anglade P.; Mistou M.-Y.

CORPORATE SOURCE: Dr. M.-Y. Mistou, Inst. Natl. de la Rech. Agronomique,

Unite Biochim./Struct. des Proteines, 78352 Jouy-en-Josas

Cedex, France. mistou@jouy.inra.fr

SOURCE: Proteomics, (1 Mar 2003) 3/3 (337-354).

Refs: 54

ISSN: 1615-9853 CODEN: PROTC7

COUNTRY: Germany

DOCUMENT TYPE: Journal; Article FILE SEGMENT: 004 Microbiology

LANGUAGE: English SUMMARY LANGUAGE: English

Lactococcus lactis is a Gram-positive bacteria, which belongs to the group of lactic acid bacteria among which several genera play an essential role in the manufacture of food products. Cytosolic proteins of L. lactis IL1403 cultivated in M17 broth have been resolved by two-dimensional gel electrophoresis using two pH gradients (pH 4-7, 4.5-5.5). More than 230 spots were identified by peptide mass fingerprints, corresponding to 25% of the predicted acid proteome. The present study made it possible to describe at the proteome level a significant number of cellular pathways (glycolysis, fermentation, nucleotide metabolism, proteolysis, fatty acid and peptidoglycan synthesis) related to important physiological processes and technological properties. It also indicated that the fermentative metabolism, which characterizes L. lactis is associated with a high expression of qlycolytic enzymes. Thirty-four proteins were matched to open reading frames for which there is no assigned function. The comparison at the proteome level of two strains of L. lactis showed an important protein polymorphism. The comparison of the proteomes of glucose- and lactose-grown cells revealed an unexpected link between the nature of the carbon source and the metabolism of pyrimidine nucleotides.

L14 ANSWER 7 OF 8 SCISEARCH COPYRIGHT 2003 THOMSON ISI on STN

ACCESSION NUMBER: 2003:286319 SCISEARCH

THE GENUINE ARTICLE: 658AT

TITLE: Proteomic analysis of Lactococcus lactis, a

lactic acid bacterium

AUTHOR: Guillot A; Gitton C; Anglade P; Mistou M Y (Reprint)

CORPORATE SOURCE: Inst Natl Rech Agron, Unite Biochim & Struct Prot, F-78352

Jouy En Josas, France (Reprint)

COUNTRY OF AUTHOR: France

SOURCE:

PROTEOMICS, (MAR 2003) Vol. 3, No. 3, pp. 337-354. Publisher: WILEY-V C H VERLAG GMBH, PO BOX 10 11 61,

D-69451 WEINHEIM, GERMANY.

ISSN: 1615-9853. Article; Journal

DOCUMENT TYPE: LANGUAGE:

English

REFERENCE COUNT:

54

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

Lactococcus lactis is a Gram-positive bacteria, which belongs to the AB group of lactic acid bacteria among which several genera play an essential role in the manufacture of food products. Cytosolic proteins of L. lactis IL1403 cultivated in M17 broth have been resolved by two-dimensional gel electrophoresis using two pH gradients (pH 4-7, 4.5-5.5). More than 230 spots were identified by peptide mass fingerprints, corresponding to 25% of the predicted acid proteome. The

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L14 ANSWER 8 OF 8 SCISEARCH COPYRIGHT 2003 THOMSON ISI on STN

ACCESSION NUMBER: 2001:251427 SCISEARCH

THE GENUINE ARTICLE: 413VM

TITLE: Genomic and proteomic analysis of microbial

function in the gastrointestinal tract of ruminants -

Review

AUTHOR: White B A (Reprint); Morrison M

CORPORATE SOURCE: Univ Illinois, Dept Anim Sci, 1207 W Gregory Dr, Urbana,

IL 61801 USA (Reprint); Univ Illinois, Dept Anim Sci, Urbana, IL 61801 USA; Univ Illinois, Dept Vet Pathobiol, Urbana, IL 61801 USA; Univ Illinois, Div Nutrit Sci,

Urbana, IL 61801 USA

COUNTRY OF AUTHOR: USA

SOURCE: ASIAN-AUSTRALASIAN JOURNAL OF ANIMAL SCIENCES, (JUN 2001)

Vol. 14, No. 6, pp. 880-884.

Publisher: ASIAN-AUSTRALASIAN ASSOC ANIMAL PRODUCTION SOCIETIES, COLLEGE AGRICULTURE LIFE SCIENCES, DEPT ANIMAL

SCIENCE TECHNOLOGY, SUWON 441-744, SOUTH KOREA.

ISSN: 1011-2367. Article; Journal

DOCUMENT TYPE: Article;
LANGUAGE: English

REFERENCE COUNT: 29

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

AΒ Rumen microbiology research has undergone several evolutionary steps: the isolation and nutritional characterization of readily cultivated microbes; followed by the cloning and sequence analysis of individual genes relevant to key digestive processes; through to the use of small subunit ribosomal RNA (SSU rRNA) sequences for cultivation-independent examination of microbial diversity. Our knowledge of rumen microbiology has expanded as a result, but the translation of this information into productive alterations of ruminal function has been rather limited. For instance, the cloning and characterization of cellulase genes in Escherichia coli has yielded some valuable information about this complex enzyme system in ruminal bacteria. SSU rRNA analyses have also confirmed that a considerable amount of the microbial diversity in the rumen is not represented in existing culture collections. However, we still have little idea of whether the key, and potentially rate-limiting, gene products and (or) microbial interactions have been identified. Technologies allowing high throughput nucleotide and protein sequence analysis have led to the emergence of two new fields of investigation, genomics and proteomics. Both disciplines can be further subdivided into functional and comparative lines of investigation. The massive accumulation of microbial DNA and protein sequence data, including complete genome sequences, is revolutionizing the way we examine microbial physiology and diversity. We describe here some examples of our use of genomics- and proteomics-based methods, to analyze the cellulase system of Ruminococcus flavefaciens FD-1 and explore the genome of Ruminococcus albus 8. At Illinois, we are using bacterial artificial chromosome (BAC) vectors to create libraries containing large (>75

kbases), contiguous segments of DNA from R. flavefaciens FD-1. Considering that every bacterium is nor a candidate for whole genome sequencing, BAC libraries offer an attractive, alternative method to perform physical and functional analyses of a bacterium's genome. Our first plan is to use these BAC clones to determine whether or not cellulases and accessory genes in R flavefaciens exist in clusters of orthologous genes (COGs). Proteomics is also being used to complement the BAC library/DNA sequencing approach. Proteins differentially expressed in response to carbon source are being identified by 2-D SDS-PAGE, followed by in-gel-digests and peptide mass mapping by MALDI-TOF Mass Spectrometry, as well as peptide sequencing by Edman degradation. At Ohio State, we have used a combination of functional proteomics, mutational analysis and differential display RT-PCR to obtain evidence suggesting that in addition to a cellulosome-like mechanism, R albus 8 possesses other mechanisms for adhesion to plant surfaces. Genome walking on either side of these differentially expressed transcripts has also resulted in two interesting observations: i) a relatively large number of genes with no matches in the current databases and; ii) the identification of genes with a high level of sequence identity to those identified, until now, in the archaebacteria. Genomics and proteomics will also accelerate our understanding of microbial interactions, and allow a greater degree of in situ analyses in the future. The challenge is to utilize genomics and proteomics to improve our fundamental understanding of microbial physiology, diversity and ecology, and overcome constraints to ruminal function.

=> s 113 and cell

L15 1282 L13 AND CELL

=> s 115 and schubert

L16 0 L15 AND SCHUBERT

=> s 115 and heterogeneous

L17 15 L15 AND HETEROGENEOUS

=> d 117 ibib abs 1-15

L17 ANSWER 1 OF 15 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN

ACCESSION NUMBER: DOCUMENT NUMBER:

2003:439217 BIOSIS PREV200300439217

TITLE:

Proteomic analysis of human vessels: Application

to atherosclerotic plaques.

AUTHOR(S):

Duran, Mari Carmen; Mas, Sebastian; Martin-Ventura, Jose

Luis; Meilhac, Olivier; Michel, Jean Baptiste;

Gallego-Delgado, Julio; Lazaro, Alberto; Tunon, Jose;

Egido, Jesus; Vivanco, Fernando (1)

CORPORATE SOURCE:

(1) Departamento de Inmunologia, Fundacion Jimenez Diaz,

Avda. Reyes Catolicos 2, 28040, Madrid, Spain:

fvivanco@fjd.es Spain

SOURCE:

Proteomics, (June 2003, 2003) Vol. 3, No. 6, pp. 973-978.

print.

ISSN: 1615-9853.

DOCUMENT TYPE:

LANGUAGE:

Article English

AB Atherosclerosis is a chronic disease that affects medium and large arteries. This process originates from the interaction between cells of the arterial wall, lipoproteins and inflammatory cells, leading to the development of complex lesions or plaques that protrude into the arterial lumen. Plaque rupture and thrombosis result in acute clinical complications such as myocardial infarction and stroke. Owing to the heterogeneous cellular composition of the

plaques, a **proteomic** analysis of the whole lesion is not appropriate. Therefore, we have studied the proteins secreted by human carotid atherosclerotic plaques, obtained by endarterectomy. Normal artery segments and different regions of the surgical pieces (noncomplicated plaque, complicated plaque with thrombus) were cultured in protein-free medium and the secreted proteins (supernatants) analyzed by two-dimensional gel electrophoresis. Normal artery segments secreted a moderate number of proteins (42 spots). However in the two-dimensional (2-D) gels (pH 3-10) of segments bearing a plaque, the number of spots increased markedly (154). The number of spots also increased (202) in the 2-D gels of artery segments with a ruptured plaque and thrombus. Thus, the more complicated the lesion, the higher the number of secreted proteins, suggesting the production of specific proteins relating to the complexity of the atherosclerotic lesion.

L17 ANSWER 2 OF 15 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN

ACCESSION NUMBER: 2002:561129 BIOSIS DOCUMENT NUMBER: PREV200200561129

TITLE: Orderly pattern of development of the autoantibody response

in (New Zealand WhiteXBXSB) F1 lupus mice: Characterization of target antigens and antigen spreading by two-dimensional

gel electrophoresis and mass spectrometry.

AUTHOR(S): Thebault, Sandrine; Gilbert, Daniele; Hubert, Marie;

Drouot, Laurent; Machour, Nadine; Lange, Catherine;

Charlionet, Roland; Tron, Francois (1)

CORPORATE SOURCE: (1) Faculte de Medecine et de Pharmacie, Institut de la

Sante et de la Recherche Medicale Unite 519, 22 Boulevard Gambetta, 76183, Rouen Cedex: françois.tron@chu-rouen.fr

France

SOURCE: Journal of Immunology, (October 1, 2002) Vol. 169, No. 7,

pp. 4046-4053. http://www.jimmunol.org/. print.

ISSN: 0022-1767.

DOCUMENT TYPE: Article LANGUAGE: English

AB

Immunoblots of a two-dimensional PAGE-separated HL-60 cell proteomic map and mass spectrometry were combined to characterize proteins targeted by autoantibodies produced by male (New Zealand WhiteXBXSB)F1 (WB) mice that develop lupus and anti-phospholipid syndrome. Analysis of sera sequentially obtained from seven individual mice at different ages showed that six proteins, vimentin, heat shock protein 60, UV excision-repair protein RAD23, alpha-enolase, heterogeneous nuclear ribonucleoprotein L, and nucleophosmin, were the targets of the B cell autoimmune response, and that autoantibodies to them were synthesized sequentially in an orderly pattern that recurred in all the male WB mice analyzed: anti-vimentin first and anti-nucleophosmin last, with anti-RAD23 and anti-heat shock protein 60, then anti-alpha-enolase and anti-heterogeneous nuclear ribonucleoprotein L Abs occuring concomitantly. Anti-vimentin reactivity always appeared before anti-cardiolipin and anti-DNA Abs, suggesting that vimentin is the immunogen initiating the autoimmune process. The pattern of HL-60 proteins recognized by female WB sera differed from that of male sera, indicating that the Y chromosome-linked autoimmune acceleration gene is not an accelerator but a strong modifier of the autoimmune response. Thus, 1) combining two-dimensional PAGE and mass spectrometry constitutes a powerful tool to identify the set of Aqs bound by autoantibodies present in a single serum and the whole autoantibody pattern of an autoimmune disease; 2) the diversification of the autoimmune response in male WB mice occurs in a predetermined pattern consistent with Ag spreading, and thus provides a useful model to further our understanding of the development of the autoantibody response in lupus.

ACCESSION NUMBER:

2003:530595 CAPLUS

DOCUMENT NUMBER:

139:193854

TITLE:

Proteomic analysis of human vessels:

Application to atherosclerotic plaques

AUTHOR(S):

Duran, Mari Carmen; Mas, Sebastian; Martin-Ventura, Jose Luis; Meilhac, Olivier; Michel, Jean Baptiste; Gallego-Delgado, Julio; Lazaro, Alberto; Tunon, Jose;

Egido, Jesus; Vivanco, Fernando

CORPORATE SOURCE:

Department Immunology, Fundacion Jimenez Diaz, Madrid,

Spain

SOURCE:

Proteomics (2003), 3(6), 973-978 CODEN: PROTC7; ISSN: 1615-9853 Wiley-VCH Verlag GmbH & Co. KGaA

PUBLISHER:

Journal

DOCUMENT TYPE: LANGUAGE:

English

Atherosclerosis is a chronic disease that affects medium and large AB arteries. This process originates from the interaction between cells of the arterial wall, lipoproteins and inflammatory cells, leading to the development of complex lesions or plagues that protrude into the arterial lumen. Plaque rupture and thrombosis result in acute clin. complications such as myocardial infarction and stroke. Owing to the heterogeneous cellular compn. of the plaques, a proteomic anal. of the whole lesion is not appropriate. Therefore, we have studied the proteins secreted by human carotid atherosclerotic plaques, obtained by endarterectomy. artery segments and different regions of the surgical pieces (noncomplicated plaque, complicated plaque with thrombus) were cultured in protein-free medium and the secreted proteins (supernatants) analyzed by two-dimensional gel electrophoresis. Normal artery segments secreted a moderate no. of proteins (42 spots). However in the two-dimensional (2-D) gels (pH 3-10) of segments bearing a plaque, the no. of spots increased markedly (154). The no. of spots also increased (202) in the 2-D gels of artery segments with a ruptured plaque and thrombus. Thus, the more complicated the lesion, the higher the no. of secreted proteins, suggesting the prodn. of specific proteins relating to the complexity of

REFERENCE COUNT:

THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L17 ANSWER 4 OF 15 CAPLUS COPYRIGHT 2003 ACS on STN

18

the atherosclerotic lesion.

ACCESSION NUMBER:

2003:3180 CAPLUS

DOCUMENT NUMBER:

138:366172

TITLE:

Proteomic analysis of the cellular proteins

induced by adaptive concentrations of hydrogen

peroxide in human U937 cells

AUTHOR(S):

Seong, Je Kyung; Kim, Do Kyun; Choi, Kun Ho; Oh, Seung Hyun; Kim, Kil Soo; Lee, Seung-Sook; Um, Hong-Duck

CORPORATE SOURCE:

Laboratory of Developmental Biology College of

Veterinary Medicine, Seoul National University, Seoul,

151-742, S. Korea

SOURCE:

Experimental and Molecular Medicine (2002), 34(5),

374-378

CODEN: EMMEF3; ISSN: 1226-3613

PUBLISHER:

Korean Society of Medical Biochemistry and Molecular

Biology

Journal

DOCUMENT TYPE: LANGUAGE: English

When cells are first exposed to low levels of oxidative stress, they develop a resistance to a subsequent challenge of the same stress, even at higher levels. Although some protein(s) induced by oxidative stress likely mediated this adaptive response, the nature of these proteins is unknown. In this study, the total proteins extd. from human U937 leukemia cells exposed to 50 .mu.M H2O2 for 24 h to induce an optimal protective response were analyzed by two-dimensional PAGE. H2O2 treatment induced elevation of level of 34 protein spots. An anal. of these spots by a matrix assocd. laser desorption/ionization time-of-flight mass spectrometry identified 28 of the H2O2-induced proteins. These include proteins involved in energy metab., translation and RNA processing, chaperoning or mediating protein folding, cellular signaling, and redox regulation, as well as a mitochondrial channel component, and an actin-bundling protein. Therefore, it appears that the cellular adaptation to oxidative stress is a complex process, and is accompanied by a modulation of diverse cellular functions.

REFERENCE COUNT: 14 THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L17 ANSWER 5 OF 15 CAPLUS COPYRIGHT 2003 ACS on STN

ACCESSION NUMBER: 2002:753691 CAPLUS

DOCUMENT NUMBER: 137:231236

TITLE: Orderly pattern of development of the autoantibody

response in (New Zealand White .times. BXSB)F1 lupus mice: characterization of target antigens and antigen spreading by two-dimensional gel electrophoresis and

mass spectrometry

AUTHOR(S): Thebault, Sandrine; Gilbert, Daniele; Hubert, Marie;

Drouot, Laurent; Machour, Nadine; Lange, Catherine;

Charlionet, Roland; Tron, Francois

CORPORATE SOURCE: Institut de la Sante et de la Recherche Medicale Unite

519, Faculte de Medecine et de Pharmacie, Hopital

Charles-Nicolle, Rouen, 76183, Fr.

SOURCE: Journal of Immunology (2002), 169(7), 4046-4053

CODEN: JOIMA3; ISSN: 0022-1767

PUBLISHER: American Association of Immunologists

DOCUMENT TYPE: Journal LANGUAGE: English

AB Immunoblots of a two-dimensional PAGE-sepd. HL-60 cell

proteomic map and mass spectrometry were combined to characterize proteins targeted by autoantibodies produced by male (New Zealand White .times. BXSB)F1 (WB) mice that develop lupus and anti-phospholipid syndrome. Anal. of sera sequentially obtained from seven individual mice at different ages showed that six proteins, vimentin, heat shock protein 60, UV excision-repair protein RAD23, .alpha.-enolase,

60, UV excision-repair protein RAD23, .alpha.-enolase, heterogeneous nuclear ribonucleoprotein L, and nucleophosmin, were the targets of the B cell autoimmune response, and that autoantibodies to them were synthesized sequentially in an orderly pattern that recurred in all the male WB mice analyzed: anti-vimentin first and anti-nucleophosmin last, with anti-RAD23 and anti-heat shock protein 60, then anti-lalpha.-enolase and anti-heterogeneous nuclear ribonucleoprotein L Abs occurring concomitantly. Anti-vimentin reactivity always appeared before anti-cardiolipin and anti-DNA Abs, suggesting that vimentin is the immunogen initiating the autoimmune process. The pattern of HL-60 proteins recognized by female WB sera differed from that of male sera, indicating that the Y chromosome-linked autoimmune acceleration gene is not an accelerator but a strong modifier of the autoimmune response. Thus, 1) combining two-dimensional PAGE and mass spectrometry constitutes a powerful tool to identify the set of Ags bound by autoantibodies present in a single serum and the whole autoantibody pattern of an autoimmune disease; 2) the diversification of the autoimmune response in male WB mice occurs in a predetd. pattern consistent with Aq spreading, and thus provides a useful model to further our understanding of the development of the autoantibody response in lupus.

REFERENCE COUNT: 51 THERE ARE 51 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L17 ANSWER 6 OF 15 MEDLINE on STN

IN-PROCESS ACCESSION NUMBER: 2003420473

PubMed ID: 12959633 DOCUMENT NUMBER: 22840803

Biomarkers, validation and pharmacokinetic-pharmacodynamic TITLE:

modelling.

AUTHOR: Colburn Wayne; Lee Jean

CORPORATE SOURCE: MDS Pharma Services, Phoenix, Arizona, USA.

CLINICAL PHARMACOKINETICS, (2003) 42 (12) 997-1022. SOURCE:

Journal code: 7606849. ISSN: 0312-5963.

PUB. COUNTRY: New Zealand

Journal; Article; (JOURNAL ARTICLE) DOCUMENT TYPE:

LANGUAGE: English

IN-PROCESS; NONINDEXED; Priority Journals FILE SEGMENT:

Entered STN: 20030909 ENTRY DATE:

Last Updated on STN: 20030909

Four elements are crucial to successful pharmacokinetic-pharmacodynamic AΒ (PK/PD) modelling and simulation for efficient and effective rational drug development: (i) mechanism-based biomarker selection and correlation to clinical endpoints; (ii) quantification of drug and/or metabolites in biological fluids under good laboratory practices (GLP); (iii) GLP-like biomarker method validation and measurements and; (iv) mechanism-based PK/PD modelling and validation. Biomarkers can provide great predictive value in early drug development if they reflect the mechanism of action for the intervention even if they do not become surrogate endpoints. PK/PD modelling and simulation can play a critical role in this process. Data from genomic and proteomics differentiating healthy versus disease states lead to biomarker discovery and identification. Multiple genes control complex diseases via hosts of gene products in biometabolic pathways and cell/organ signal transduction. Pilot exploratory studies should be conducted to identify pivotal biomarkers to be used for predictive clinical assessment of disease progression and the effect of drug intervention. Most biomarkers are endogenous macromolecules, which could be measured in biological fluids. Many exist in heterogeneous forms with varying activity and immunoreactivity, posting challenges for bioanalysis. Reliable and selective assays could be validated under a GLP-like environment for quantitative methods. While the need for consistent reference standards and quality control monitoring during sample analysis for biomarker assays are similar to that of drug molecules, many biomarkers have special requirements for sample collection that demand a wellcoordinated team management. Bioanalytical methods should be validated to meet study objectives at various drug development stages, and possess adequate performance to quantify biochemical responses specific to the target disease progression and drug intervention. Protocol design to produce sufficient data for PK/PD modelling would be more complex than that of PK. Knowledge of mechanism from discovery and preclinical studies are helpful for planning clinical study designs in cascade, sequential, crossover or replicate mode. The appropriate combination of biomarker identification and selection, bioanalytical methods development and validation for drugs and biomarkers, and mechanism-based PK/PD models for fitting data and predicting future clinical endpoints/outcomes provide powerful insights and guidance for effective and efficient rational drug development, toward

L17 ANSWER 7 OF 15 MEDLINE on STN

IN-PROCESS ACCESSION NUMBER: 2003305117 DOCUMENT NUMBER: 22717145 PubMed ID: 12833522

TITLE: Proteomic analysis of human vessels: application

safe and efficacious medicine for individual patients.

to atherosclerotic plaques.

AUTHOR: Duran Mari Carmen; Mas Sebastian; Martin-Ventura Jose Luis;

> Meilhac Olivier; Michel Jean Baptiste; Gallego-Delgado Julio; Lazaro Alberto; Tunon Jose; Egido Jesus; Vivanco

Fernando

CORPORATE SOURCE: Department Immunology, Fundacion Jimenez Diaz, Madrid,

Spain.

Proteomics, (2003 Jun) 3 (6) 973-8. SOURCE:

Journal code: 101092707. ISSN: 1615-9853. Germany: Germany, Federal Republic of

PUB. COUNTRY: DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: IN-PROCESS; NONINDEXED; Priority Journals

ENTRY DATE: Entered STN: 20030701

Last Updated on STN: 20030718

Atherosclerosis is a chronic disease that affects medium and large AB arteries. This process originates from the interaction between cells of the arterial wall, lipoproteins and inflammatory cells, leading to the development of complex lesions or plagues that protrude into the arterial lumen. Plaque rupture and thrombosis result in acute clinical complications such as myocardial infarction and stroke. Owing to the heterogeneous cellular composition of the plaques, a proteomic analysis of the whole lesion is not appropriate. Therefore, we have studied the proteins secreted by human carotid atherosclerotic plaques, obtained by endarterectomy. artery segments and different regions of the surgical pieces (noncomplicated plaque, complicated plaque with thrombus) were cultured in protein-free medium and the secreted proteins (supernatants) analyzed by two-dimensional gel electrophoresis. Normal artery segments secreted a moderate number of proteins (42 spots). However in the two-dimensional (2-D) gels (pH 3-10) of segments bearing a plaque, the number of spots increased markedly (154). The number of spots also increased (202) in the 2-D gels of artery segments with a ruptured plaque and thrombus. the more complicated the lesion, the higher the number of secreted proteins, suggesting the production of specific proteins relating to the complexity of the atherosclerotic lesion.

L17 ANSWER 8 OF 15 MEDLINE on STN

ACCESSION NUMBER: 2002489889 MEDLINE

DOCUMENT NUMBER: 22229570 PubMed ID: 12244208

Orderly pattern of development of the autoantibody response TITLE:

in (New Zealand White x BXSB) F1 lupus mice:

characterization of target antiqens and antiqen spreading

by two-dimensional gel electrophoresis and mass

spectrometry.

AUTHOR: Thebault Sandrine; Gilbert Daniele; Hubert Marie; Drouot

Laurent; Machour Nadine; Lange Catherine; Charlionet

Roland; Tron Francois

CORPORATE SOURCE: Institut de la Sante et de la Recherche Medicale Unite 519,

Faculte de Medecine et de Pharmacie, Hopital

Charles-Nicolle, Rouen, France.

SOURCE: JOURNAL OF IMMUNOLOGY, (2002 Oct 1) 169 (7) 4046-53.

Journal code: 2985117R. ISSN: 0022-1767.

PUB. COUNTRY:

United States Journal; Article; (JOURNAL ARTICLE) DOCUMENT TYPE:

LANGUAGE:

FILE SEGMENT: Abridged Index Medicus Journals; Priority Journals

ENTRY MONTH: 200211

ENTRY DATE: Entered STN: 20020928

> Last Updated on STN: 20021213 Entered Medline: 20021112

Immunoblots of a two-dimensional PAGE-separated HL-60 cell proteomic map and mass spectrometry were combined to characterize proteins targeted by autoantibodies produced by male (New Zealand White x BXSB)F(1) (WB) mice that develop lupus and anti-phospholipid syndrome. Analysis of sera sequentially obtained from seven individual mice at

different ages showed that six proteins, vimentin, heat shock protein 60, UV excision-repair protein RAD23, alpha-enolase, heterogeneous nuclear ribonucleoprotein L, and nucleophosmin, were the targets of the B cell autoimmune response, and that autoantibodies to them were synthesized sequentially in an orderly pattern that recurred in all the male WB mice analyzed: anti-vimentin first and anti-nucleophosmin last, with anti-RAD23 and anti-heat shock protein 60, then anti-alpha-enolase and anti-heterogeneous nuclear ribonucleoprotein L Abs occuring concomitantly. Anti-vimentin reactivity always appeared before anti-cardiolipin and anti-DNA Abs, suggesting that vimentin is the immunogen initiating the autoimmune process. The pattern of HL-60 proteins recognized by female WB sera differed from that of male sera, indicating that the Y chromosome-linked autoimmune acceleration gene is not an accelerator but a strong modifier of the autoimmune response. Thus, 1) combining two-dimensional PAGE and mass spectrometry constitutes a powerful tool to identify the set of Ags bound by autoantibodies present in a single serum and the whole autoantibody pattern of an autoimmune disease; 2) the diversification of the autoimmune response in male WB mice occurs in a predetermined pattern consistent with Ag spreading, and thus provides a useful model to further our understanding of the development of the autoantibody response in lupus.

L17 ANSWER 9 OF 15 MEDLINE on STN ACCESSION NUMBER: 1999127506 MEDLINE

DOCUMENT NUMBER: 99127506 PubMed ID: 9928545

TITLE: Experimental pathology and breast cancer genetics: new

technologies.

AUTHOR: Osin P; Shipley J; Lu Y J; Crook T; Gusterson B A

CORPORATE SOURCE: Section of Cell Biology and Experimental Pathology,

Institute of Cancer Research, Haddow Laboratories, Sutton,

Surrey, UK.

SOURCE: RECENT RESULTS IN CANCER RESEARCH, (1998) 152 35-48. Ref:

37

Journal code: 0044671. ISSN: 0080-0015. PUB. COUNTRY: GERMANY: Germany, Federal Republic of

PUB. COUNTRY: GERMANY: Germany, Federal Republic of DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

General Review; (REVIEW)

(REVIEW, TUTORIAL)

LANGUAGE: English

Engitsii

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199903

ENTRY DATE: Entered STN: 19990326

Last Updated on STN: 19990326 Entered Medline: 19990312

AΒ The goal is to understand the critical events in tumour development and to apply this understanding to new approaches to diagnosis, prevention and treatment. It is clear that breast cancer is a heterogeneous disease at the molecular level, raising the possibility of a future functional classification based on mechanisms rather than morphology. These molecular phenotypes will also confer predictive value on the potential of the tumour to invade, metastasise and respond to or resist new therapeutic strategies. Studies of the genome in individuals are predicted also to enable the identification of polymorphisms that are associated with increased susceptibility to environmental factors, in addition to possibly explaining de novo variations in responses to drugs and radiation. The difficulty is how to identify which, of the approximately 30,000 genes expressed by a typical cancer cell alone or in combination, are the ones involved in these processes The majority of breast cancers have such a multitude of molecular changes that it is difficult to distinguish between those that are critical to tumour progression and those that are epiphenomena of genetic instability and abnormalities in DNA repair. The identification of the

earliest events in carcinogenesis must be the best hope, as it will then be possible to target the events that predispose to other secondary changes before they occur. Genomics and **proteomics** is the current hope to take us forward. This involves the application of a number of new technologies to facilitate the profiling of individual tumours, including laser-guided microdissection of microscopic lesions, comparative genomic hybridisation and loss of heterozygosity analysis of DNA using microarray technology to study DNA and expressed RNAs and protein profiling using 2D gel mass spectroscopy. With over 100,000 mRNAs and proteins to examine in complex tissues and in various combinations, there is obviously going to be a requirement for a large investment in computing power (bioinformatics) to facilitate the analysis of these data in relation to the clinical characteristics of the individual tumour and the patient.

L17 ANSWER 10 OF 15 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED.

on STN

ACCESSION NUMBER: 2003270566 EMBASE

TITLE: Proteomic analysis of human vessels: Application

to atherosclerotic plaques.

AUTHOR: Duran M.C.; Mas S.; Martin-Ventura J.L.; Meilhac O.; Michel

J.B.; Gallego-Delgado J.; Lazaro A.; Tunon J.; Egido J.;

Vivanco F.

CORPORATE SOURCE: Dr. F. Vivanco, Departamento de Inmunologia, Fundacion

Jimenez Diaz, Avda. Reyes Catolicos, 2-28040-Madrid, Spain.

fvivanco@fjd.es

SOURCE: Proteomics, (1 Jun 2003) 3/6 (973-978).

Refs: 18

ISSN: 1615-9853 CODEN: PROTC7

COUNTRY: Germany

DOCUMENT TYPE: Journal; Conference Article

FILE SEGMENT: 005 General Pathology and Pathological Anatomy

018 Cardiovascular Diseases and Cardiovascular Surgery

029 Clinical Biochemistry

LANGUAGE: English SUMMARY LANGUAGE: English

Atherosclerosis is a chronic disease that affects medium and large arteries. This process originates from the interaction between cells of the arterial wall, lipoproteins and inflammatory cells, leading to the development of complex lesions or plaques that protrude into the arterial lumen. Plaque rupture and thrombosis result in acute clinical complications such as myocardial infarction and stroke. Owing to the heterogeneous cellular composition of the plaques, a proteomic analysis of the whole lesion is not appropriate. Therefore, we have studied the proteins secreted by human carotid atherosclerotic plaques, obtained by endarterectomy. Normal artery segments and different regions of the surgical pieces (noncomplicated plaque, complicated plaque with thrombus) were cultured in protein-free medium and the secreted proteins (supernatants) analyzed by two-dimensional gel electrophoresis. Normal artery segments secreted a moderate number of proteins (42 spots). However in the two-dimensional (2-D) gels (pH 3-10) of segments bearing a plaque, the number of spots increased markedly (154). The number of spots also increased (202) in the 2-D gels of artery segments with a ruptured plaque and thrombus. Thus, the more complicated the lesion, the higher the number of secreted proteins, suggesting the production of specific proteins relating to the complexity of the atherosclerotic lesion.

L17 ANSWER 11 OF 15 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED. on STN

ACCESSION NUMBER: 2003130598 EMBASE

TITLE: Utility of peptide-protein affinity complexes in

proteomics: Identification of interaction partners of a tumor suppressor p21([141-160])(cip1/waf1) peptide.

Gururaja T.L.; Li W.; Payan D.G.; Anderson D.C. AUTHOR:

CORPORATE SOURCE: Dr. T.L. Gururaja, Rigel Pharmaceuticals Inc., 240 East

Grand Avenue, South San Francisco, CA 94080, United States.

tgururaja@rigel.com

SOURCE: Journal of Peptide Research, (1 Apr 2003) 61/4 (163-176).

Refs: 65

ISSN: 1397-002X CODEN: JPERFA

COUNTRY: United Kingdom DOCUMENT TYPE: Journal; Article

FILE SEGMENT: 027 Biophysics, Bioengineering and Medical

Instrumentation

029 Clinical Biochemistry

LANGUAGE: English SUMMARY LANGUAGE: English

We used a N-biotinylated peptide analog of the C-terminal domain of the tumor suppressor protein, p21(cip1/waf1) to elucidate peptide/protein interacting partners. The C-terminal domain of p21(cip1/waf1) protein spanning 141-160 amino acid residues is known to bind PCNA and this interaction is important in many biological processes including cell-cycle control. This C-terminal 20-mer efficiently extracts PCNA in the presence of a variety of N- or C-terminally attached affinity tags. Using difference silver stained 2D gels combined with in-gel tryptic digests, we identified the difference spots using MALDI-TOF mass spectrometry-based peptide mass fingerprinting followed by a database search using PROFOUND against NCBIs human nonredundant protein sequence data bank. Identified spots include the p48 subunit of chromatin assembly factor-I, the heat shock 70 protein analog BiP, calmodulin, nucleolin and a spot similar in size to dimeric PCNA. In contrast, microcapillary ion-trap LC-MS/MS analysis of a tryptic digest of entire affinity extracts derived from both control and experimental runs followed by database searches using SEQUEST confirmed the presence of most of the above proteins. This strategy also identified hnRNPA1, HPSP90.alpha., HSP40 and T-complex protein 1, a protein similar to prothymosin, and a possible allelic variant of the p21(cip1/waf1) protein. The use of N-biotinylated peptide derived from the C-terminal domain of p21(cip1/waf1) protein in proteomic analysis exemplified here suggests that peptides obtained from intracellular functional screens could also potentially

serve as efficient baits to discover new drug targets.

L17 ANSWER 12 OF 15 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED. on STN

ACCESSION NUMBER: 2002339598 EMBASE

TITLE: Orderly pattern of development of the autoantibody response

in (New Zealand White x BXSB)F(1) lupus mice:

Characterization of target antigens and antigen spreading

by two-dimensional gel electrophoresis and mass

spectrometry.

AUTHOR: Thebault S.; Gilbert D.; Hubert M.; Drouot L.; Machour N.;

Lange C.; Charlionet R.; Tron F.

Dr. F. Tron, Inst. Sante/Rech. Medicale Unite 519, Faculte CORPORATE SOURCE:

de Medecine et de Pharmacie, 22 boulevard Gambetta, 76183

Rouen Cedex, France. francois.tron@chu-rouen.fr

SOURCE: Journal of Immunology, (1 Oct 2002) 169/7 (4046-4053).

Refs: 51

ISSN: 0022-1767 CODEN: JOIMA3

COUNTRY: United States DOCUMENT TYPE: Journal; Article

FILE SEGMENT: General Pathology and Pathological Anatomy 005

026 Immunology, Serology and Transplantation

LANGUAGE: English



Immunoblots of a two-dimensional PAGE-separated HL-60 cell proteomic map and mass spectrometry were combined to characterize proteins targeted by autoantibodies produced by male (New Zealand White x BXSB)F(1) (WB) mice that develop lupus and anti-phospholipid syndrome. Analysis of sera sequentially obtained from seven individual mice at different ages showed that six proteins, vimentin, heat shock protein 60, UV excision-repair protein RAD23, .alpha.-enolase, heterogeneous nuclear ribonucleoprotein L, and nucleophosmin, were the targets of the B cell autoimmune response, and that autoantibodies to them were synthesized sequentially in an orderly pattern that recurred in all the male WB mice analyzed: anti-vimentin first and anti-nucleophosmin last, with anti-RAD23 and anti-heat shock protein 60, then anti-.alpha.-enolase and anti-heterogeneous nuclear ribonucleoprotein L Abs occuring concomitantly. Anti-vimentin reactivity always appeared before anti-cardiolipin and anti-DNA Abs, suggesting that vimentin is the immunogen initiating the autoimmune process. The pattern of HL-60 proteins recognized by female WB sera differed from that of male sera, indicating that the Y chromosome-linked autoimmune acceleration gene is not an accelerator but a strong modifier of the autoimmune response. Thus, 1) combining two-dimensional PAGE and mass spectrometry constitutes a powerful tool to identify the set of Ags bound by autoantibodies present in a single serum and the whole autoantibody pattern of an autoimmune disease; 2) the diversification of the autoimmune response in male WB mice occurs in a predetermined pattern consistent with Ag spreading, and thus provides a useful model to further our understanding of the development of the auto-antibody response in lupus.

L17 ANSWER 13 OF 15 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED.

on STN

ACCESSION NUMBER:

2002337436 EMBASE

TITLE:

Tissue microdissection and its applications in pathology.

AUTHOR:

Curran S.; Murray G.I.

CORPORATE SOURCE:

G.I. Murray, Department of Pathology, University of

Aberdeen, Foresterhill, Aberdeen AB25 2ZD, United Kingdom.

g.i.murray@abdn.ac.uk

SOURCE:

COUNTRY:

Current Diagnostic Pathology, (2002) 8/3 (183-192).

Refs: 42

ISSN: 0968-6053 CODEN: CDPAFN

DOCUMENT TYPE:

United Kingdom

FILE SEGMENT:

Journal; General Review

005 General Pathology and Pathological Anatomy

016

LANGUAGE:

English

SUMMARY LANGUAGE: English

Human tissues are composed of complex admixtures of different cell types. In order to examine a specific population of cells, it is necessary to isolate them from surrounding cells: this process can be problematic. Tissue microdissection may be defined as a process of isolating a morphologically distinct population of cells from a tissue section or cytological preparation composed of a mixture of heterogeneous cell types. The purpose of tissue microdissection is to provide a homogeneous sample of cells, in a form suitable for further analysis. Microdissection is becomingly increasingly important, due to the development of a variety of powerful molecular analytical techniques, which must be applied to samples of a high degree of purity/homogeneity in order to provide clinically and biologically meaningful results. A variety of different methods of microdissection have been developed. This review describes the basic principles of tissue microdissection techniques, and considers their applications in the field of human pathology. .COPYRGT. 2002 Elsevier Science Ltd. All rights reserved.

L17 ANSWER 14 OF 15 SCISEARCH COPYRIGHT 2003 THOMSON ISI on STN

2003:595589 SCISEARCH ACCESSION NUMBER:

THE GENUINE ARTICLE: 697CL

Proteomic analysis of human vessels: Application TITLE:

to atherosclerotic plaques

Duran M C; Mas S; Martin-Ventura J L; Meilhac O; Michel J AUTHOR:

B; Gallego-Delgado J; Lazaro A; Tunon J; Egido J; Vivanco

CORPORATE SOURCE: Fdn Jimenez Diaz, Dept Immunol, Avda Reyes Catollicos 2,

> Madrid 28040, Spain (Reprint); Fdn Jimenez Diaz, Dept Immunol, Madrid 28040, Spain; Autonomous Univ Madrid, Renal & Vasc Lab, Fdn Jimenez Diaz, E-28049 Madrid, Spain; CHU Xavier Bichat, Unit 460 INSERM, Paris, France; Fdn Jimenez Diaz, Dept Cardiol, E-28040 Madrid, Spain; Univ Complutense, Preteom Unit, Dept Biochem & Mol Biol 1,

E-28040 Madrid, Spain

COUNTRY OF AUTHOR:

Spain; France

SOURCE:

PROTEOMICS, (JUN 2003) Vol. 3, No. 6, pp. 973-978. Publisher: WILEY-V C H VERLAG GMBH, PO BOX 10 11 61,

D-69451 WEINHEIM, GERMANY.

ISSN: 1615-9853. Article; Journal

DOCUMENT TYPE: LANGUAGE:

English

REFERENCE COUNT:

18

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

AB Atherosclerosis is a chronic disease that affects medium and large arteries. This process originates from the interaction between cells of the arterial wall, lipoproteins and inflammatory cells, leading to the development of complex lesions or plaques that protrude into the arterial lumen. Plaque rupture and thrombosis result in acute clinical complications such as myocardial infarction and stroke. Owing to the heterogeneous cellular composition of the plaques, a proteomic analysis of the whole lesion is not appropriate. Therefore, we have studied the proteins secreted by human carotid atherosclerotic plaques, obtained by endarterectomy. Normal artery segments and different regions of the surgical pieces (noncomplicated plaque, complicated plaque with thrombus) were cultured in protein-free medium and the secreted proteins (supernatants) analyzed by two-dimensional gel electrophoresis. Normal artery segments secreted a moderate number of proteins (42 spots). However in the two-dimensional (2-D) gels (pH 3-10) of segments bearing a plaque, the number of spots increased markedly (154). The number of spots also increased (202) in the 2-D gels of artery segments with a ruptured plaque and thrombus. Thus, the more complicated the lesion, the higher the number of secreted proteins, suggesting the production of specific proteins relating to the complexity of the atherosclerotic lesion.

L17 ANSWER 15 OF 15 SCISEARCH COPYRIGHT 2003 THOMSON ISI on STN

ACCESSION NUMBER: 2002:815911 SCISEARCH

THE GENUINE ARTICLE: 598QV

TITLE:

Orderly pattern of development of the autoantibody response in (New Zealand White x BXSB) F-1 lupus mice:

Characterization of target antigens and antigen spreading by two-dimensional gel electrophoresis and mass

spectrometry

AUTHOR:

Thebault S; Gilbert D; Hubert M; Drouot L; Machour N;

Lange C; Charlionet R; Tron F (Reprint)

CORPORATE SOURCE:

Inst Sante & Rech, Med Unite 519, Fac Med & Pharm, 22 Blvd Gambetta, F-76183 Rouen, France (Reprint); Hop Charles Nicolle, Inst Sante & Rech, Med Unit 519, Fac Med & Pharm, Rouen, France; Inst Rech Chim Organ Fine, Lab Spect, Inst

Fed Rech Multidisciplinaires Peptides, Rouen, France

COUNTRY OF AUTHOR:

France

SOURCE:

JOURNAL OF IMMUNOLOGY, (1 OCT 2002) Vol. 169, No. 7, pp.

4046-4053.

Publisher: AMER ASSOC IMMUNOLOGISTS, 9650 ROCKVILLE PIKE,

BETHESDA, MD 20814 USA.

ISSN: 0022-1767.

DOCUMENT TYPE:

Article; Journal

LANGUAGE:

English

REFERENCE COUNT:

51

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS AΒ Immunoblots of a two-dimensional PAGE-separated HL-60 cell proteomic map and mass spectrometry were combined to characterize proteins targeted by autoantibodies produced by male (New Zealand White X BXSB)F-1 (WB) mice that develop lupus and anti-phospholipid syndrome. Analysis of sera sequentially obtained from seven individual mice at different ages showed that six proteins, vimentin, heat shock protein 60, UV excision-repair protein RAD23, alpha-enolase, heterogeneous nuclear ribonucleoprotein L, and nucleophosmin, were the targets of the B cell autoimmune response, and that autoantibodies to them were synthesized sequentially in an orderly pattern that recurred in all the male WB mice analyzed: anti-vimentin first and anti-nucleophosmin last, with anti-RAD23 and anti-heat shock protein 60, then anti-alpha-enolase and anti-heterogeneous nuclear ribonucleoprotein L Abs occuring concomitantly. Anti-vimentin reactivity always appeared before anti-cardiolipin and anti-DNA Abs, suggesting that vimentin is the immunogen initiating the autoimmune process. The pattern of HL-60 proteins recognized by female WB sera differed from that of male sera, indicating that the Y chromosome-linked autoimmune acceleration gene is not an accelerator but a strong modifier of the autoimmune response. Thus, 1) combining two-dimensional PAGE and mass spectrometry constitutes a powerful tool to identify the set of Aqs bound by autoantibodies present in a single serum and the whole autoantibody pattern of an autoimmune disease; 2) the diversification of the autoimmune response in male WB mice occurs in a predetermined pattern consistent with Ag spreading, and thus provides a useful model to further our understanding of the development of the autoantibody response in lupus.

=> s 115 and protein chips

L18 1 L15 AND PROTEIN CHIPS

=> d 118 abs

L18 ANSWER 1 OF 1 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN

AN 2003-175091 [17] WPIDS

CR 2003-183869 [18]

AB WO 200294454 A UPAB: 20030317

NOVELTY - Fabrication of microdevices involves a vapor deposition coating process, where the coating includes functional groups having an intrinsic reactivity to react with target molecules.

DETAILED DESCRIPTION - An INDEPENDENT CLAIM is included for a microdevice article for separation of biomolecules in an electrical field comprising a polymeric coating for capturing molecules at the surface, which bind at least a part of the biomolecules subjected to screening, only temporarily allowing their subsequent release.

USE - For parallel analysis of biomolecules (particularly hydrogels, polyelectrolytes or temp-sensitive molecules) and for separation of biomolecules in an electrical field (all claimed). The biomolecules include biologically active molecules in pharmaceutical technology such as in the fields of drug discovery, **proteomics**, genomics, high-throughput screening, clinical diagnostics, and other applications

such as manufacturing of cell arrays, immobilization of drugs for tissue engineering, microreactors, surfaces for protein and DNA screening and on electro-optical devices e.g. miniaturized analytical systems, biomedical devices, tools for chemistry and biochemistry and systems for fundamental research; such as useful for decoltation of implantable devices e.g. heart valves, pacemakers, stents, embolization coils, bone substitution, hip substitution, bone screws, vascular grafts etc.; improved scaffold for tissue engineering, plates for in vitro cell culture, resins for protein synthesis, microchip based diagnostic screening, protein purification, DNA purification, DNA chips, protein chips, arrays of quantum dots, electro-optical devices, and coating of three-dimensional structures such as membranes, micro-reactors, micro-channels, foams, scaffold etc. The functionalized coating can be used for microdevices made of different materials such as polymers, composites, silicon, semiconductors, glass or metal.

ADVANTAGE - The process is a simple one-step process to prepare high-content and high-throughput screening surfaces and can be quickly scaled-up. The polymer interfaces are highly reactive and contain chemical groups having intrinsic reactivity to react with target molecules, in which, at least part of capturing molecules specifically bind to biomolecules that are subject to screening, in binding pairs such as antibody/antigen, antibody/hapten, enzyme/substrate, integrin/extracellular matrix component, biomolecule/cell, cell/cell, carrier protein/substrate, lectine/carbohydrate, protein/carbohydrate, carbohydrate/carbohydrate, cell adhesion molecule/cell surface receptor, receptor/hormone, receptor/cytokine, protein/DNA, protein/RNA, peptide/DNA, two DNA single strains, DNA/RNA, DNA/DNA, in which either of both partners of the couples serve as capturing molecules. The use of capturing molecules avoids the step of surface modification of the bulk material. The coating process provides an increased surface concentration of functional groups with a defined and controlled ratio, compared to conventional methods such as plasma treatment. Due to chemically stable background of the deposited polymer, aging effects as a consequence of interactions with analyte solutions can be reduced or ruled out. Due to the mild character of the deposition process, side reactions are suppressed and the deposited films are homogenous with respect to their chemical structure and topology. Gradients in the functionalized groups in the coatings can also be produced by establishment of temperature gradients at the substrate subjected to coating. Straightforward synthesis achieves spatially directed immobilization of biomolecules. The once deposited film can be subjected to further modification using conventional methods. The coating step is substrate independent, thus provides a generic approach to microstructuring of microdevices. The method provides unique characteristics of various instantly reacting functional groups, a defined and chemically stable polymer base layer with defined properties, non-degradability, the possibility to create activity gradients, and the feasibility for patterning. The method overcomes the restrictions associated with gold/alkanethiolates-based techniques such as surface modification of the bulk material and side reactions including the fabrication or incorporation of potently harmful chemicals prior to immobilization, and previous deposition of gold, the method maintains the intrinsic advantages of soft lithography e.g. accuracy, broad availability and low costs. The manufacture of the microdevices by soft lithography, allows easy scale-up by replication, involving replica molding comprising casting of prepolymer against a master and preparing a negative replica of the master in polydiemethylsiloxane The coating can be produced in any shape ranging from three dimensional to porous structure. Dwg.0/0

L18 ANSWER 1 OF 1 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN

AN 2003-175091 [17] WPIDS

CR 2003-183869 [18]

AB WO 200294454 A UPAB: 20030317

NOVELTY - Fabrication of microdevices involves a vapor deposition coating **process**, where the coating includes functional groups having an intrinsic reactivity to react with target molecules.

DETAILED DESCRIPTION - An INDEPENDENT CLAIM is included for a microdevice article for separation of biomolecules in an electrical field comprising a polymeric coating for capturing molecules at the surface, which bind at least a part of the biomolecules subjected to screening, only temporarily allowing their subsequent release.

USE - For parallel analysis of biomolecules (particularly hydrogels, polyelectrolytes or temp-sensitive molecules) and for separation of biomolecules in an electrical field (all claimed). The biomolecules include biologically active molecules in pharmaceutical technology such as in the fields of drug discovery, proteomics, genomics, high-throughput screening, clinical diagnostics, and other applications such as manufacturing of cell arrays, immobilization of drugs for tissue engineering, microreactors, surfaces for protein and DNA screening and on electro-optical devices e.g. miniaturized analytical systems, biomedical devices, tools for chemistry and biochemistry and systems for fundamental research; such as useful for decoltation of implantable devices e.g. heart valves, pacemakers, stents, embolization coils, bone substitution, hip substitution, bone screws, vascular grafts etc.; improved scaffold for tissue engineering, plates for in vitro cell culture, resins for protein synthesis, microchip based diagnostic screening, protein purification, DNA purification, DNA chips, protein chips, arrays of quantum dots, electro-optical devices, and coating of three-dimensional structures such as membranes, micro-reactors, micro-channels, foams, scaffold etc. The functionalized coating can be used for microdevices made of different materials such as polymers, composites, silicon, semiconductors, glass or metal.

ADVANTAGE - The process is a simple one-step process to prepare high-content and high-throughput screening surfaces and can be quickly scaled-up. The polymer interfaces are highly reactive and contain chemical groups having intrinsic reactivity to react with target molecules, in which, at least part of capturing molecules specifically bind to biomolecules that are subject to screening, in binding pairs such as antibody/antigen, antibody/hapten, enzyme/substrate, integrin/extracellular matrix component, biomolecule/cell, cell/cell, carrier protein/substrate, lectine/carbohydrate, protein/carbohydrate, carbohydrate/carbohydrate, cell adhesion molecule/cell surface receptor, receptor/hormone, receptor/cytokine, protein/DNA, protein/RNA, peptide/DNA, two DNA single strains, DNA/RNA, DNA/DNA, in which either of both partners of the couples serve as capturing molecules. The use of capturing molecules avoids the step of surface modification of the bulk material. The coating process provides an increased surface concentration of functional groups with a defined and controlled ratio, compared to conventional methods such as plasma treatment. Due to chemically stable background of the deposited polymer, aging effects as a consequence of interactions with analyte solutions can be reduced or ruled out. Due to the mild character of the deposition process, side reactions are suppressed and the deposited films are homogenous with respect to their chemical structure and topology. Gradients in the functionalized groups in the coatings can also be produced by establishment of temperature gradients at the substrate subjected to coating. Straightforward synthesis achieves spatially directed immobilization of biomolecules. The once deposited film can be subjected

to further modification using conventional methods. The coating step is substrate independent, thus provides a generic approach to microstructuring of microdevices. The method provides unique characteristics of various instantly reacting functional groups, a defined and chemically stable polymer base layer with defined properties, non-degradability, the possibility to create activity gradients, and the feasibility for patterning. The method overcomes the restrictions associated with gold/alkanethiolates-based techniques such as surface modification of the bulk material and side reactions including the fabrication or incorporation of potently harmful chemicals prior to immobilization, and previous deposition of gold, the method maintains the intrinsic advantages of soft lithography e.g. accuracy, broad availability and low costs. The manufacture of the microdevices by soft lithography, allows easy scale-up by replication, involving replica molding comprising casting of prepolymer against a master and preparing a negative replica of the master in polydiemethylsiloxane The coating can be produced in any shape ranging from three dimensional to porous structure.

Dwg.0/0

ACCESSION NUMBER: 2003-175091 [17] WPIDS

CROSS REFERENCE: 2003-183869 [18]
DOC. NO. NON-CPI: N2003-137926
DOC. NO. CPI: C2003-045709

TITLE: Fabrication of microdevices for parallel analysis of

biomolecules involves a vapor deposition coating **process** with the coating including functional groups having an intrinsic reactivity to react with

target molecules.

DERWENT CLASS: A26 A89 B04 D16 M13 P42 S03

INVENTOR(S): LAHANN, J

PATENT ASSIGNEE(S): (LAHA-I) LAHANN J

COUNTRY COUNT: 100

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2002094454 A1 20021128 (200317) * EN 16

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TR TZ UG ZM ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EC EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ OM PH PL PT RO RU SD SE SG SI SK SL TJ TM TN TR TT TZ UA UG US UZ VN YU ZA ZM ZW

DE 10124873 A1 20021205 (200317)

APPLICATION DETAILS:

PATENT NO KI	[ND	API	PLICATION	DATE
WO 2002094454	A1	WO	2002-US16326	20020522
DE 10124873	A1	DE	2001-10124873	20010522

PRIORITY APPLN. INFO: DE 2001-10125872 20010522; DE 2001-10124873 20010522

=> s 115 and automatic?

L19 4 L15 AND AUTOMATIC?

=> d l19 abs ibib 1-4

L19 ANSWER 1 OF 4 CAPLUS COPYRIGHT 2003 ACS on STN

The objective of this paper is to present a methodol. for developing and AΒ calibrating models of complex reaction/transport systems. In particular, the complex network of biochem. reaction/transport processes and their spatial organization make the development of a predictive model of a living cell a grand challenge for the 21st century. However, advances in reaction/transport modeling and the exponentially growing databases of genomic, proteomic, metabolic, and bioelec. data make cell modeling feasible, if these two elements can be automatically integrated in an unbiased fashion. In this paper, we present a procedure to integrate data with a new cell model, Karyote, that accounts for many of the phys. processes needed to attain the goal of predictive modeling. Our integration methodol. is based on the use of information theory. The model is integrated with a variety of types and qualities of exptl. data using an objective error assessment approach. Data that can be used in this approach include NMR, spectroscopy, microscopy, and elec. potentiometry. The approach is demonstrated on the well-studied Trypanosoma brucei system. A major obstacle for the development of a predictive cell model is that the complexity of these systems makes it unlikely that any model presently available will soon be complete in terms of the set of processes accounted for. Thus, one is faced with the challenge of calibrating and running an incomplete model. We present a probability functional method that allows the integration of exptl. data and soft information such as choice of error measure, a priori information, and phys. motivated regularization to address the incompleteness challenge.

ACCESSION NUMBER:

2003:726762 CAPLUS

TITLE:

Toward Automated Cell Model Development

through Information Theory

AUTHOR(S):

Sayyed-Ahmad, A.; Tuncay, K.; Ortoleva, Peter J.

CORPORATE SOURCE: Center for Cell and Virus Theory Department of

Chemistry, Indiana University, Bloomington, IN, 47405,

USA

SOURCE:

Journal of Physical Chemistry ACS ASAP

CODEN: JPCHAX; ISSN: 0022-3654

PUBLISHER:

American Chemical Society

DOCUMENT TYPE:

Journal

LANGUAGE:

English

REFERENCE COUNT:

32 THERE ARE 32 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L19 ANSWER 2 OF 4 CAPLUS COPYRIGHT 2003 ACS on STN

When presented with a mixt. of intact proteins, electrospray ionization with Fourier-transform mass spectrometry (ESI-FTMS) has the capability to obtain direct fragmentation information from isolated ions. However, the automation of this capability has not been achieved to date. We have developed software for unattended acquisition of protein tandem mass spectrometry (MS/MS) data and batch processing of the resulting files for identification of whole proteins. Mixts. of both protein stds. (8-29 kDa) and Methanococcus jannaschii cytosolic proteins (up to six components < 20 kDa) were infused via an autosampler, and MS/MS data were acquired without human intervention. The acquisition software recognizes ESI charge state patterns, generates protein-specific isolation waveforms on-the-fly, and fragments ions using two different IR laser times. In addn. to protein stds., five wild-type proteins (7-14 kDa) were identified automatically with 100% sequence coverage from the M. jannaschii database. The software underpins a measurement platform for sample-dependent acquisition of MS/MS data for whole proteins, a crit. step to realize proteomics with 100% sequence coverage in a higher throughput setting.

ACCESSION NUMBER:

2002:785182 CAPLUS

DOCUMENT NUMBER:

138:85921



TITLE: Fourier-transform mass spectrometry for automated

fragmentation and identification of 5-20 kda proteins

in mixtures

AUTHOR(S): Johnson, Jeffrey R.; Meng, Fanyu; Forbes, Andrew J.;

Cargile, Benjamin J.; Kelleher, Neil L.

CORPORATE SOURCE: Department of Chemistry, University of Illinois,

Urbana, IL, USA

SOURCE: Electrophoresis (2002), 23(18), 3217-3223

> CODEN: ELCTDN; ISSN: 0173-0835 Wiley-VCH Verlag GmbH & Co. KGaA

PUBLISHER: DOCUMENT TYPE: Journal

English LANGUAGE:

THERE ARE 38 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 38 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L19 ANSWER 3 OF 4 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN

2003-371684 [35] WPIDS AN

AΒ WO2003011553 A UPAB: 20030603

> NOVELTY - Preparation of an artificial cell membrane (A) by dispensing a membrane lipid (I) across an aperture then thinning (I) by applying suction at the sides.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for:

- (1) a method for producing a membrane (A') in which a liquid polymer is used in place of (I) and heat is applied to (II) before, during or after some stage in the process; and
- (2) an apparatus for preparing (A) or (A') comprising a member with at least one aperture and at least one adjacent side channel for applying

USE - (A), including bilayer lipid membranes, are useful e.g. in ion-channel sensing; DNA sensors; proteomics and design/synthesis of proteins. Also arrays of (A) on a chip are useful for classifying and assaying agents for their effects on cellular receptors and to improve characterization of pathogens. Polymeric membranes prepared similarly are useful as fluid filters.

ADVANTAGE - The method is suitable for automated production of membranes that retain stability and integrity without a cellular infrastructure, without requiring cumbersome techniques such as the patch clamp method. The status of the membrane can be monitored continuously during use and any damaged membranes can be regenerated automatically. When used in conjunction with in vitro translation/transcription systems, complex membrane-bound receptor systems can be formed, optionally with reagents drawn, electrically, to specific sites, allowing control over which proteins are expressed form a large array. This makes possible simultaneous analysis of many complex molecular combinations without interference from extraneous components (as happens in cellular systems).

DESCRIPTION OF DRAWING(S) - Schematic diagram of the apparatus for preparing membranes, and the stages involved in lipid membrane formation (BLM = bilayer lipid membrane).

Dwg.3/34

ACCESSION NUMBER: 2003-371684 [35] WPIDS

DOC. NO. CPI: C2003-098539

TITLE: Preparation of artificial cell or polymeric

membranes, useful e.g. as DNA sensors, prepared by applying suction to a solution across an opening.

DERWENT CLASS: A89 B04 D16

INVENTOR(S): PARAMESWARAN, L; YOUNG, A

PATENT ASSIGNEE(S): (MASI) MASSACHUSETTS INST TECHNOLOGY

COUNTRY COUNT: 100

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LΑ PG WO 2003011553 A1 20030213 (200335) * EN 50

RW: AT BE BG CH CY CZ DE DK EA EE ES FI FR GB GH GM GR IE IT KE LS LU

MC MW MZ NL OA PT SD SE SK SL SZ TR TZ UG ZM ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EC EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ OM PH PL PT RO RU SD SE SG SI SK SL TJ TM TN TR TT TZ UA UG US UZ VN YU ZA ZM

US 2003062657 A1 20030403 (200335)

APPLICATION DETAILS:

P	ATENT	NO K	IND		API	PLICATION	DATE
		3011553 3062657		Provisional	US	2002-US24266 2001-309259P 2002-209321	20020730 20010731 20020730

PRIORITY APPLN. INFO: US 2001-309259P 20010731; US 2002-209321 20020730

L19 ANSWER 4 OF 4 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED. on STN

AΒ Two-dimensional electrophoresis is a widely used method for separating a large number of proteins from complex protein mixtures and for revealing differential patterns of protein expressions. In the computer-assisted proteome research, the comparison of protein separation profiles involves several heuristic steps, ranging from protein spot detection to matching of unknown spots. An important prerequisite for efficient protein spot matching is the image warping step, where the geometric relationship between the gel profiles is modeled on the basis of a given set of known corresponding spots, so-called landmarks, and the locations of unknown spots are predicted using the optimized model. Traditionally, polynomial functions together with least squares optimization has been used, even though this approach is known to be incapable of modeling all the complex distortions inherent in electrophoretic data. To satisfy the need of more flexible gel distortion correction, a hierarchical grid transformation method with stochastic optimization is presented. The method provides an adaptive multiresolution model between the gels, and good correction performance in the practical crossvalidation tests suggests that automatic warping of gel images could be based on this approach. We believe that the proposed model also has significance in the ultimate comparison of corresponding protein spots since the matching

process should benefit from the closeness of the true spot pairs.

ACCESSION NUMBER: 2002442751 EMBASE

TITLE: Hierarchical grid transformation for image warping in the

analysis of two-dimensional electrophoresis gels.

AUTHOR: Salmi J.; Aittokallio T.; Westerholm J.; Griese M.;

Rosengren A.; Nyman T.A.; Lahesmaa R.; Nevalainen O. Dr. T. Aittokallio, Turku Centre for Computer Science,

Lemminkaisenkatu 14A, FIN-20520 Turku, Finland.

tero.aittokallio@utu.fi

SOURCE: Proteomics, (1 Nov 2002) 2/11 (1504-1515).

Refs: 21

ISSN: 1615-9853 CODEN: PROTC7

COUNTRY: Germany

CORPORATE SOURCE:

DOCUMENT TYPE: Journal; Article

FILE SEGMENT: 029 Clinical Biochemistry

LANGUAGE: English SUMMARY LANGUAGE: English

=> FIL STNGUIDE

SINCE FILE COST IN U.S. DOLLARS TOTAL ENTRY SESSION 118.55 FULL ESTIMATED COST 163.36

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FILE CONTAINS CURRENT INFORMATION. LAST RELOADED: Sep 26, 2003 (20030926/UP).

=> FIL BIOSIS, CAPLUS, MEDLINE, WPIDS, EMBASE, SCISEARCH

COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION 0.42 FULL ESTIMATED COST 163.78

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION CA SUBSCRIBER PRICE 0.00 -5.86

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FILE 'SCISEARCH' ENTERED AT 16:25:49 ON 29 SEP 2003 COPYRIGHT 2003 THOMSON ISI

=> d hist

L3

(FILE 'HOME' ENTERED AT 15:45:30 ON 29 SEP 2003)

FILE 'BIOSIS, CAPLUS, MEDLINE, WPIDS, EMBASE, SCISEARCH' ENTERED AT 15:45:38 ON 29 SEP 2003

E SCHUBERT

E E3 AND WALTER

E SCHUBERT

1 S E3 AND WALTER 1.1

1 S E3 AND WALTER L2

229 S E3 AND CELL 5 S L3 AND TARGET L4

FILE 'STNGUIDE' ENTERED AT 15:52:18 ON 29 SEP 2003

O S CELL SPECIFIC TARGET AND PROCESS L5 O S CELL SPECIFIC TARGET L6 0 S CELL- SPECIFIC TARGET L7 L8O S CELL- SPECIFIC AND SCHUBERT L9 0 S MELK L10 4 S PROTEOMICS FILE 'BIOSIS, CAPLUS, MEDLINE, WPIDS, EMBASE, SCISEARCH' ENTERED AT 16:06:49 ON 29 SEP 2003 FILE 'STNGUIDE' ENTERED AT 16:07:16 ON 29 SEP 2003 FILE 'BIOSIS, CAPLUS, MEDLINE, WPIDS, EMBASE, SCISEARCH' ENTERED AT 16:07:32 ON 29 SEP 2003 L11 16796 S PROTEOMIC? L12 1 S L11 AND MELK L13 2438 S PROTEOMIC AND PROCESS 8 S L13 AND CULTIVATED L14L15 1282 S L13 AND CELL 0 S L15 AND SCHUBERT L16 L17 15 S L15 AND HETEROGENEOUS L18 1 S L15 AND PROTEIN CHIPS L19 4 S L15 AND AUTOMATIC? FILE 'STNGUIDE' ENTERED AT 16:21:29 ON 29 SEP 2003 FILE 'BIOSIS, CAPLUS, MEDLINE, WPIDS, EMBASE, SCISEARCH' ENTERED AT 16:25:49 ON 29 SEP 2003 => s 115 and endotheli? L20 27 L15 AND ENDOTHELI? => dup rem 120 PROCESSING COMPLETED FOR L20 16 DUP REM L20 (11 DUPLICATES REMOVED) => d 121 1-16 abs ibib L21 ANSWER 1 OF 16 CAPLUS COPYRIGHT 2003 ACS on STN Genomic actions and/or proteomic interactions for pathophysiol. processes and for physiol. processes are detd. at assocd. redox state conditions. Protein interactions are correlated with oxygen tensions. Identification of markers for disease including epitopes is effected in the presence of simulated redox state perturbations.

Screening for previously unknown receptors and activating ligands is

carried out in the presence of alteration of redox state. ACCESSION NUMBER: 2003:300520 CAPLUS

DOCUMENT NUMBER: 138:283690

TITLE: Determination of **proteomic** interaction and

genomic action in the presence of associated redox state conditions and physiological, pathophysiological

and drug screening applications

INVENTOR(S): Stamler, Jonathan S.

PATENT ASSIGNEE(S): USA

SOURCE: U.S. Pat. Appl. Publ., 9 pp.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

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US 2001-977693
                                                            20011016
    US 2003073088
                      Α1
                            20030417
                                           WO 2002-US31571 20021015
    WO 2003033721
                      A2
                            20030424
                            20030814
    WO 2003033721
                      Α3
            AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
             PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ,
            UA, UG, UZ, VC, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU,
            TJ, TM
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG,
            CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
             PT, SE, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR,
            NE, SN, TD, TG
PRIORITY APPLN. INFO.:
                                        US 2001-977693
                                                         A 20011016
```

ANSWER 2 OF 16 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN DUPLICATE 1

AB Atherosclerosis is a progressive and complex pathophysiological process occurring in large arteries. Although it is of multifactorial origin, the disease develops at preferential sites along the vasculature in regions experiencing specific hemodynamic conditions that are predisposed to endothelial dysfunction. The exact mechanisms allowing endothelial cells to discriminate between plaque-free and plaque-prone flows remain to be explored. To investigate such mechanisms, we performed a proteomic analysis on endothelial cells exposed in vitro to these two-flow patterns. A few spots on the two-dimensional gel had an intensity that was differentially regulated by plaque-free versus plaque-prone flows. One of them was further investigated and identified as macrophage-capping protein (Cap G), a member of the gelsolin protein superfamily. A 2-fold increase of Cap G protein and a 5-fold increase of Cap G mRNA were observed in cells exposed to a plaque-free flow as compared with static cultures. This increase was not observed in cells exposed to plaque-prone flow. Plaque-free flow induced a corresponding increase in nuclear and cytoskeletal-associated Cap G. Finally, overexpression of Cap G in transfection assays increased the motility potential of endothelial cells. These observations together with the known functions of Cap G suggest that Cap G may contribute to the protective effect exerted by plaque-free flow on endothelial cells. On the contrary, in cells exposed to a plaque-prone flow, no induction of Cap G expression could be

observed. ACCESSION NUMBER: 2003:444600 BIOSIS DOCUMENT NUMBER: PREV200300444600

TITLE: Cap G, a gelsolin family protein modulating protective

effects of unidirectional shear stress.

AUTHOR(S): Pellieux, Corinne; Desgeorges, Alain; Pigeon, Christelle

Haziza; Chambaz, Celine; Yin, Helen; Hayoz, Daniel;

Silacci, Paolo (1)

(1) Division of Hypertension and Vascular Medicine, Centre CORPORATE SOURCE:

Hospitalier Universitaire Vaudois, 1015, Lausanne,

Switzerland: paolo.silacci@epfl.ch Switzerland

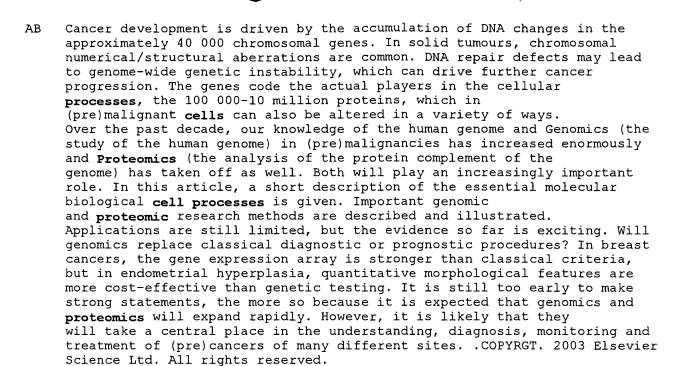
Journal of Biological Chemistry, (August 1 2003) Vol. 278, SOURCE:

No. 31, pp. 29136-29144. print.

ISSN: 0021-9258.

DOCUMENT TYPE: Article LANGUAGE: English

L21 ANSWER 3 OF 16 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED. on STN



ACCESSION NUMBER:

2003201384 EMBASE

TITLE:

Genomics and proteomics in cancer.

AUTHOR:

Baak J.P.A.; Path F.R.C.; Hermsen M.A.J.A.; Meijer G.;

Schmidt J.; Janssen E.A.M.

CORPORATE SOURCE:

J.P.A. Baak, Department of Pathology, Central Hospital for Rogaland, Box 8001, 4068 Stavanger, Norway. baja@sir.no

SOURCE:

European Journal of Cancer, (2003) 39/9 (1199-1215).

Refs: 112

ISSN: 0959-8049 CODEN: EJCAEL

COUNTRY:

United Kingdom

DOCUMENT TYPE:

Journal; General Review

FILE SEGMENT:

016 Cancer

029 Clinical Biochemistry

LANGUAGE:

English

SUMMARY LANGUAGE: English

L21 ANSWER 4 OF 16 SCISEARCH COPYRIGHT 2003 THOMSON ISI on STN

AB Blood platelets are important components of haemostasis. After their

activation they cause healing of wounds by forming plugs and initiate repair processes. One important event in regulating this activation is the phosphorylation/dephosphorylation of multiple proteins on various tyrosine, serine and threonine residues. To understand the exact molecular mechanisms in platelet activation it is essential to identify proteins involved in the signalling pathways and to localise and characterise their phosphorylation sites. After treatment with P-32 and separation by 2D-PAGE using different pI ranges, phosphorylated platelet proteins were detected by autoradiography. Phosphotyrosine-containing proteins were assigned by immunoblotting with an anti-phosphotyrosine antibody. Another approach for the identification of phosphorylated proteins was immunoprecipitation of tyrosine-phosphorylated proteins using an anti-phosphotyrosine antibody. Protein spots/bands of interest were excised from the get, digested with trypsin and analysed by MALDI-TOF-MS and nano-LC-ESI-MS/MS, respectively. Several phosphorylated proteins could be identified and the localisation of some in vivo phosphorylation sites was possible.

ACCESSION NUMBER:

2003:772136 SCISEARCH

THE GENUINE ARTICLE: 718PL

TITLE: D

Differential analysis of phosphorylated proteins in

resting and thrombin-stimulated human platelets

AUTHOR: Marcus K (Reprint); Moebius J; Meyer H E

CORPORATE SOURCE: Ruhr Univ Bochum, Med Proteom Ctr, ZKF 141, Univ Str 150,

D-44780 Bochum, Germany (Reprint); Ruhr Univ Bochum, Med

Proteom Ctr, ZKF 141, D-44780 Bochum, Germany

COUNTRY OF AUTHOR:

Germany

SOURCE:

ANALYTICAL AND BIOANALYTICAL CHEMISTRY, (AUG 2003) Vol.

376, No. 7, pp. 973-993.

Publisher: SPRINGER-VERLAG BERLIN, HEIDELBERGER PLATZ 3,

D-14197 BERLIN, GERMANY.

ISSN: 1618-2642.

DOCUMENT TYPE:

Article; Journal

LANGUAGE:

English

REFERENCE COUNT:

96

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

L21 ANSWER 5 OF 16 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN DUPLICATE 2

The endothelium is a single layer of cells lining the AΒ inside face of all blood vessels. It constitutes a major metabolic organ which is critically involved in the generation and the regulation of multiple physiological and pathological processes such as coagulation, hemostasis, inflammation, atherosclerosis, angiogenesis and cancerous metastasis dissemination. In order to increase our knowledge about the protein content and the main biological pathways of human vascular endothelial cells, we have undertaken the proteomic analysis of the most explored present endothelial cell model, i.e. primo-cultures of human umbilical vein endothelial cells (HUVECs). Using low levels of protein loads (apprx30 mug), the association of two-dimensional electrophoresis with matrix-assisted laser desorption/ionization-time of flight mass spectrometry, liquid chromatography-tandem mass spectrometry and database interrogations allowed us to identify 53 proteins of suspected endothelial origin in quiescent HUVECs. Beside cytoskeletal proteins such as actin, tubulin, tropomyosin and vimentin, we identified various proteins more especially implicated in cellular motility and plasticity (e.g. cofilin, F-actin capping protein and prefoldin), in regulation of apoptosis and senescence (protease inhibitor 9, glucose related proteins, heat shock proteins, thioredoxin peroxidase, nucleophosmin) as well as other proteins implicated in coagulation (annexin V, high mobility group protein), antigen presentation (valosin containing protein and ubiquitin carboxyl terminal hydrolase isozyme L1) and enzymatic capabilities (glutathione-S-transferase, protein disulfide isomerases, lactate deshydrogenase).

ACCESSION NUMBER: 2003:329994 BIOSIS DOCUMENT NUMBER: PREV200300329994

TITLE: Proteomic study of human umbilical vein

endothelial cells in culture.

AUTHOR(S): Bruneel, Arnaud (1); Labas, Valerie; Mailloux, Agnes;

Sharma, Sanjiv; Vinh, Joelle; Vaubourdolle, Michel; Baudin,

Bruno

CORPORATE SOURCE: (1) Service de Biochimie A, Hopital Saint-Antoine, AP-HP,

184 Rue du Faubourg Saint-Antoine, 75571, Paris Cedex 12,

France: arnaud.bruneel@sat.ap-hop-paris.fr France

SOURCE: Proteomics, (May 2003, 2003) Vol. 3, No. 5, pp. 714-723.

print.

ISSN: 1615-9853.

DOCUMENT TYPE:

Article

LANGUAGE:

English

L21 ANSWER 6 OF 16 SCISEARCH COPYRIGHT 2003 THOMSON ISI on STN

AB Homeostasis of the intracellular ionic concentration, in particular

that of hydrogen ions, is pivotal to the maintenance of cell function and viability. Nonetheless, pH fluctuations in both the intracellular and the extracellular compartments can occurr during development, in physiological processes and in disease. The influence of pH variations on gene expression has been studied in different model systems, but only for a limited number of genes. We have performed a broad range analysis of the patterns of gene expression in normal human dermal fibroblasts at two different pH values (in the presence and in the absence of serum), with the aim of getting a deeper insight into the regulation of the transcriptional program as a response to a pH change. Using the Affymetrix gene chip system, we found that the expression of 2068 genes (out of 12 565) was modulated by more than two-fold at 24, 48 or 72 h after the shift of the culture medium pH to a more acidic value, stanniocalcin 1 being a remarkable example of a strongly up-regulated gene. Genes displaying a modulated pattern of expression included, among others, cell cycle regulators (consistent with the observation that acidic pH abolishes the growth of fibroblasts in culture) and relevant extracellular matrix (ECM) components. Extracellular matrix protein 2, a protein with a restricted pattern of expression in adult human tissues, was found to be remarkably overexpressed as a consequence of serum starvation. Since ECM components, whose expression is controlled by pH, have been used as targets for biomolecular intervention, we have complemented the Affymetrix analysis with a two-dimensional polyacrylamide gel electrophoresis analysis of proteins which are differentially secreted by fibroblasts at acidic or basic pH. Mass spectrometric analysis of more than 650 protein spots allowed the identification of 170 protein isoforms or fragments, belonging to 40 different proteins. Some proteins were only expressed at basic pH (including, for instance, tetranectin), while others (e.g., agrin) were only detectable at acidic pH. Some of the identified proteins may represent promising candidate targets for biomedical applications, e.g., for antibody-mediated vascular targeting strategies.

ACCESSION NUMBER: 2003:465783 SCISEARCH

THE GENUINE ARTICLE: 679TA

TITLE. Made

TITLE: Modulation of gene expression by extracellular pH

variations in human fibroblasts: A transcriptomic and

proteomic study

AUTHOR: Bumke M A; Neri D (Reprint); Elia G

CORPORATE SOURCE: Swiss Fed Inst Technol, Inst Pharmaceut Sci,

Winterthurerstr 190, CH-8057 Zurich, Switzerland

(Reprint); Swiss Fed Inst Technol, Inst Pharmaceut Sci,

CH-8057 Zurich, Switzerland

COUNTRY OF AUTHOR:

Switzerland

SOURCE:

PROTEOMICS, (MAY 2003) Vol. 3, No. 5, pp. 675-688. Publisher: WILEY-V C H VERLAG GMBH, PO BOX 10 11 61,

D-69451 WEINHEIM, GERMANY.

ISSN: 1615-9853.

DOCUMENT TYPE:

Article; Journal

LANGUAGE:

English

79

REFERENCE COUNT:

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

L21 ANSWER 7 OF 16 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED. on STN

AB Several new PLA(2)s have been identified based on their nucleotide gene sequences. They were classified mainly into three groups: cytosolic PLA(2) (cPLA(2)), secretary PLA(2) (sPLA(2)), and intracellular PLA(2) (iPLA(2)). They differ from each other in terms of substrate specificity, Ca(2+) requirement and lipid modification. The questions that still remain to be addressed are the subcellular localization and differential regulation of the isoforms in various cell types and under different physiological conditions. It is required to identify the downstream events

that occur upon PLA(2) activation, particularly target protein or metabolic pathway for liberated arachidonic acid or other fatty acids. Understanding the same will greatly help in the development of potent and specific pharmacological modulators that can be used for basic research and clinical applications. The information of the human and other genomes of PLA(2)s, combined with the use of proteomics and genetically manipulated mouse models of different diseases, will illuminate us about the specific and potentially overlapping roles of individual phospholipases as mediators of physiological and pathological processes. Hopefully, such understanding will enable the development of specific agents aimed at decreasing the potential contribution of individual secretary phospholipases to vascular diseases. The signaling cascades involved in the activation of cPLA(2) by mitogen activated protein kinases (MAPKs) is now evident. It has been demonstrated that p44 MAPK phosphorylates cPLA(2) and increases its activity in cells and tissues. The phosphorylation of cPLA(2) at ser505 occurs before the increase in intracellular Ca(2+) that facilitate the binding of the lipid binding domain of cPLA(2) to phospholipids, promoting its translocation to cellular membranes and AA release. Recently, a negative feed back loop for cPLA(2) activation by MAPK has been proposed. If PLA(2) activation in a given model depends on PKC, PKA, cAMP, or MAPK then inhibition of these phosphorylating enzymes may alter activities of PLA(2) isoforms during cellular injury. Understanding the signaling pathways involved in the activation/deactivation of PLA(2) during cellular injury will point to key events that can be used to prevent the cellular injury. Furthermore, to date, there is limited information available regarding the regulation of iPLA(2) or sPLA(2) by these pathways. .COPYRGT. 2002 Elsevier Science Inc. All rights reserved.

ACCESSION NUMBER: 2003192191 EMBASE

TITLE: Phospholipase A(2) isoforms: A perspective.

AUTHOR: Chakraborti S.

CORPORATE SOURCE: S. Chakraborti, Dept. of Biochemistry and Biophysics,

University of Kalyani, Kalyani 741235, West Bengal, India.

s chakraborti@hotmail.com

SOURCE: Cellular Signalling, (1 Jul 2003) 15/7 (637-665).

Refs: 325

ISSN: 0898-6568 CODEN: CESIEY

COUNTRY: United States

DOCUMENT TYPE: Journal; General Review

FILE SEGMENT: 029 Clinical Biochemistry 037 Drug Literature Index

LANGUAGE: English SUMMARY LANGUAGE: English

L21 ANSWER 8 OF 16 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED. on STN

A number of high resolution two-dimensional electrophoresis (2-DE) reference maps for bovine tissues and biological fluids have been determined for animals in basal state. Among the 1863 distinct protein features detected in samples of liver, kidney, muscle, plasma and red blood cells, 509 species were identified and associated to 209 different genes. Difficulties in the identification were related to the poorly characterized Bos taurus genome and were solved by a combined matrix-assisted laser desorption/ionisation-mass spectrometry and liquid chromatography-electrospray ionization tandem mass spectrometry approach. The experimental output allowed us to establish a 2-DE database accessible through the World Wide Web network at the URL address (http://www.iabbam.na.cnr.it/Biochem). These reference maps may serve as a tool in future veterinary medical studies aimed at the evaluation of changes in protein repertoire for altered animal physiological conditions and infectious diseases, to the definition of molecular markers for novel diagnostic kits and vaccines, as well as the characterization of protein

modifications in bovine materials following technological

processes used in the food industry. 2003156060 EMBASE ACCESSION NUMBER:

Proteins from bovine tissues and biological fluids: TITLE:

Defining a reference electrophoresis map for liver, kidney,

muscle, plasma and red blood cells.

Talamo F.; D'Ambrosio C.; Arena S.; Del Vecchio P.; Ledda AUTHOR:

L.; Zehender G.; Ferrara L.; Scaloni A.

CORPORATE SOURCE: Dr. A. Scaloni, Proteomics/Mass Spectrom. Laboratory,

I.S.P.A.A.M., National Research Council, via Argine 1085,

80147 Naples, Italy. a.scaloni@iabbam.na.cnr.it

Proteomics, (1 Apr 2003) 3/4 (440-460). SOURCE:

Refs: 40

ISSN: 1615-9853 CODEN: PROTC7

Germany COUNTRY:

DOCUMENT TYPE: Journal; Article

Clinical Biochemistry FILE SEGMENT: 029

LANGUAGE: English SUMMARY LANGUAGE: English

L21 ANSWER 9 OF 16 CAPLUS COPYRIGHT 2003 ACS on STN DUPLICATE 3

An increase in permeability of the blood-brain barrier is a crit. event in AΒ

the pathophysiol. process of multiple sclerosis and other

neurodegenerative diseases. Tumor necrosis factor .alpha. (TNF.alpha.) is

known to play a crucial role in this process and is a powerful

activator of endothelial cell inflammatory responses.

Although many reports describe effects of TNF.alpha. activation in

endothelial cells, the mol. mechanisms specific for

activation of cerebral endothelial cells remains

unclear. The objective of this study was to identify potential

pharmaceutical targets for the treatment of multiple sclerosis using mol.

profiling techniques. Gene expression measurements (Affymetrix Hu6800 oligonucleotide arrays) and proteomics (two-dimensional gel

electrophoresis and mass spectrometry) were applied to analyze early

alterations in human cerebral endothelial cells (HCEC)

activated by TNF.alpha.. Human umbilical vein endothelial

cells (HUVEC) were used as the ref. system. The results presented show that HCEC and HUVEC respond similarly with respect to cell

adhesion mols., chemotaxis, apoptosis and oxidative stress mols. However, nuclear factors NFkB1 and NFkB2, plasminogen activator inhibitor 1 and

cofilin 1 are examples of cerebral specific responses. Our results indicate involvements of the urokinase plasminogen activator system and

cytoskeletal rearrangements unique to TNF.alpha. activation of cerebral endothelial cells.

Gene and protein expression profiling of human cerebral endothelial cells

activated with tumor necrosis factor-.alpha.

2003:567392 CAPLUS

Franzen, Bo; Duvefelt, Kristina; Jonsson, Carina; AUTHOR(S):

Engelhardt, Britta; Ottervald, Jan; Wickman, Monica;

Yang, Yang; Schuppe-Koistinen, Ina

CNS & Pain Control, Molecular Sciences, Gene and CORPORATE SOURCE:

Protein Technology, Local Discovery Research Area,

AstraZeneca R&D, Soedertaelje, Swed.

SOURCE: Molecular Brain Research (2003), 115(2), 130-146

CODEN: MBREE4; ISSN: 0169-328X

PUBLISHER: Elsevier Science B.V.

DOCUMENT TYPE: Journal LANGUAGE: English

ACCESSION NUMBER:

TITLE:

L21 ANSWER 10 OF 16 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN

Tumor microenvironmental factors, possibly from angiogenic

processes, alter endothelial cell surface protein expression. Accessible to the circulation, the tumor vascular endothelium is a logical target for drug or gene delivery. Transcytotic vesicles (caveolae) on the surface of the endothelium provide a pathway for overcoming the endothelial cell barrier for delivery to underlying tumor cells. We purified luminal endothelial cell plasma membranes and its caveolae directly from normal and tumor-bearing tissues and resolved the membrane proteins by SDS-PAGE and 2D gel electrophoresis to reveal extensive heterogeneity of cell surface protein expression between tissues as well as apparent tumor-induced markers. Mass spectrometry and database searching provided key sequences of tryptic peptides of tumor-associated targets. Using antibodies to the appropriate specific polypeptides, we have confirmed the tumor-induction of the proteins. One of the tumor-induced proteins, designated TE3, is expressed in endothelial caveolae of solid tumors but not normal organs. Biodistribution analysis and whole body imaging (gamma scintigraphy) using TE3 antibody injected IV show the accessibility of TE3 with significant and selective tumor tissue accumulation in just 1 hour. Targeting the endothelium and its caveolae are worthy strategies for tumor targeting in vivo.

ACCESSION NUMBER:

2003:401903 BIOSIS

DOCUMENT NUMBER:

PREV200300401903

TITLE:

Proteomic mapping of tumor neovasculature in vivo

for specific targeting and imaging of solid tumors.

AUTHOR(S):

Oh, Phil (1); Subbiah, Krishnan; Testa, Jacqueline; Czarny, Malgorzata; Smith, Traci; Hearn, Kally; Wempren, Alexina;

Schnitzer, Melinda; Schnitzer, Jan E.

CORPORATE SOURCE:

(1) Vascular Biology, Sidney Kimmel Cancer Center, 10835 Altman Row, San Diego, CA, 92121, USA: poh@skcc.org, ksubbiah@skcc.org, jtesta@skcc.org, mczarny@skcc.org, tsmith@skcc.org, khearn@skcc.org, awempren@skcc.org,

SOURCE:

mschnitzer@skcc.org, jschnitzer@skcc.org USA FASEB Journal, (March 2003, 2003) Vol. 17, No. 4-5, pp. Abstract No. 171.9. http://www.fasebj.org/. e-file.

Meeting Info.: FASEB Meeting on Experimental Biology: Translating the Genome San Diego, CA, USA April 11-15, 2003

FASEB

. ISSN: 0892-6638.

DOCUMENT TYPE:

Conference

LANGUAGE: English

ANSWER 11 OF 16 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN

2002-657552 [70] ANWPIDS

2001-607180 [69] CR

WO 200263412 A UPAB: 20030214

NOVELTY - Observations are provided for the members of multiple objects allocated to pre-existing categories. Properties of latent classes are estimated from the distinguishable sets of latent classes associated with the members of objects and corresponding to the pre-existing categories.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are included for the following:

- (1) object classification method;
- (2) computer implemented latent class identification method;
- (3) method of identifying genes linked to cellular phenotype;
- (4) method of identifying gene linked to metastatic properties of cancer;
 - (5) method of screening a drug;
 - (6) method of identifying genes linked to a disease of interest;
 - (7) membership rule generation method;
- (8) method of identifying a gene linked to metastatic properties of tumor;

(9) method of identifying genes of known and unknown functions; and (10) object image analyzing method.

USE - For identifying latent class for analysis of large amount of information for sequence analysis, gene expression and proteomics in the field of biology and for identification of genes linked to diseases such as leukemias such as acute leukemia, acute lymphocytic leukemia, acute myelocytic leukemia, myeloblastic, promyelocytic, myelomonocytic, monocytic erythroleukemia chronic leukemia, chronic myelocytic leukemia, chronic lymphocytic leukemia, polycythemia vera, lymphoma, Hodgkin's disease, non-Hodgkin's disease, multiple myeloma, Waldenstrom's macroglobulinemia, heavy chain disease, solid tumors sarcomas, carcinomas including fibrosarcoma, myxosarcoma, liposarcoma, chondrosarcoma, osteogenic sarcoma, chordoma, angiosarcoma, endotheliosarcoma, lymphangiosarcoma, Kaposi's sarcoma, lymphangioendotheliosarcoma, synovioma, mesothelioma, Ewing's tumor, leiomyosarcoma, rhabdomyosarcoma, colon carcinoma, pancreatic cancer, breast cancer, ovarian cancer, prostate cancer, squamous cell carcinoma, basal cell carcinoma, adenocarcinoma, sweat gland carcinoma, sebaceous gland carcinoma, papillary carcinoma, papillary adenocarcinomas, cystadenocarcinoma, medullary carcinoma, bronchogenic carcinoma, renal cell carcinoma, hepatoma, bile duct carcinoma, choriocarcinoma, seminoma, embryonal carcinoma, Wilms' tumor, cervical cancer, uterine cancer, testicular tumor, lung carcinoma, small cell lung carcinoma, bladder carcinoma, epithelial carcinoma, glioma, astrocytoma, ependymoma, craniopharyngioma, medulloblastoma, pinealoma, hemangioblastoma, acoustic neuroma, oligodendroglioma, meningioma, melanoma, neuroblastoma, retinoblastoma and other types of tumors including virally induced cancers and for identification of disabilities, undesirable interactions between medications, co-morbidities, laboratory results and clinical characteristics linked to processes of aging, disease, cancer, diabetes, pregnancy or other clinical or pathological conditions in humans.

ADVANTAGE - Incorportes explanatory models with intentionally weak structural assumptions to prevent the imposition of artificial patterns on the data and thereby makes the models useful for complex data exploration. Estimates broad expression patterns over genes and over **cell** or tissue samples and allows quantitative determination of new biological knowledge. Assigns individual gene probabilities of membership in specific patterns and allows the quantity uncertainty associated with the allocating elements among sets of interpretable categories. Enables to conduct formal hypothesis testing, evaluate whether an identified gene pattern is different from the null hypothesis pattern and incorporates complex model structures used to exploit external biological knowledge.

DESCRIPTION OF DRAWING(S) - The figure shows the serum stimulation patterns including median estimates and 95% confidence intervals from analysis of serum stimulated fibroblasts, demonstrating both time-dependent increases and decreases in gene expression, as represented by positive and negative expression patterns.

Dwg.1/16

ACCESSION NUMBER: 2002-657552 [70] WPIDS

CROSS REFERENCE: 2001-607180 [69]
DOC. NO. NON-CPI: N2002-519853
DOC. NO. CPI: C2002-184535

TITLE:

Latent classes identification method for identifying genes, involves estimating properties of latent classes from distinguishable sets of latent classes associated

with members of objects allocated to pre-existing

DERWENT CLASS: B04 D16 T01
INVENTOR(S): LAZARIDIS, E

PATENT ASSIGNEE(S): (LAZA-I) LAZARIDIS E; (UYSF-N) UNIV SOUTH FLORIDA

COUNTRY COUNT: 97

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2002063412 A2 20020815 (200270) * EN 81

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ

NL OA PT SD SE SL SZ TR TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EC EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR

KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PH PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

US 2002169730 A1 20021114 (200277)

US 2003023385 A1 20030130 (200311)

APPLICATION DETAILS:

PATENT NO K	IND	API	PLICATION	DATE
WO 2002063412	A2	WO	2001-US26672	20010828
US 2002169730	A1	WO	2001-US3616	20010205
		US	2001-940878	20010829
US 2003023385	A1	WO	2001-US3616	20010205
		US	2001-913498	20010816

PRIORITY APPLN. INFO: WO 2001-US3616 20010205

L21 ANSWER 12 OF 16 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED. on STN

AΒ The prognosis of hepatocellular carcinoma (HCC) still remains dismal, although many advances in its clinical study have been made. It is important for tumor control to identify the factors that predispose patients to death. With new discoveries in cancer biology, the pathological and biological prognostic factors of HCC have been studied quite extensively. Analyzing molecular markers (biomarkers) with prognostic significance is a complementary method. A large number of molecular factors have been shown to associate with the invasiveness of HCC, and have potential prognostic significance. One important aspect is the analysis of molecular markers for the cellular malignancy phenotype. These include alterations in DNA ploidy, cellular proliferation markers (PCNA, Ki-67, Mcm2, MIB1, MIA, and CSEIL/CAS protein), nuclear morphology, the p53 gene and its related molecule MDM2, other cell cycle regulators (cyclin A, cyclin D, cyclin E, cdc2, p27, p73), oncogenes and their receptors (such as ras, c-myc, c-fms, HGF, c-met, and erb-B receptor family members), apoptosis related factors (Fas and FasL), as well as telomerase activity. Another important aspect is the analysis of molecular markers involved in the process of cancer invasion and metastasis. Adhesion molecules (E-cadherin, catenins, serum intercellular adhesion molecule-1, CD44 variants), proteinases involved in the degradation of extracellular matrix (MMP-2, MMP-9, uPA, uPAR, PAI), as well as other molecules have been regarded as biomarkers for the malignant phenotype of HCC, and are related to prognosis and therapeutic outcomes. Tumor angiogenesis is critical to both the growth and metastasis of cancers including HCC, and has drawn much attention in recent years. Many angiogenesis-related markers, such as vascular endothelial growth factor (VEGF), basic fibroblast growth factor (bFGF), platelet-derived endothelial cell growth factor (PD-ECGF), thrombospondin (TSP), angiogenin, pleitrophin, and endostatin (ES) levels, as well as intratumor microvessel density (MVD) have been evaluated and found to be of prognostic significance. Body fluid (particularly blood and urinary) testing for biomarkers is easily accessible and useful in clinical patients. The prognostic significance of

circulating DNA in plasma or serum, and its genetic alterations in HCC are other important trends. More attention should be paid to these two areas in future. As the progress of the human genome project advances, so does a clearer understanding of tumor biology, and more and more new prognostic markers with high sensitivity and specificity will be found and used in clinical assays. However, the combination of some items, i. e., the pathological features and some biomarkers mentioned above, seems to be more practical for now.

ACCESSION NUMBER: 2002230965 EMBASE

TITLE: The prognostic molecular markers in hepatocellular

carcinoma.

AUTHOR: Qin L.-X.; Tang Z.-Y.

CORPORATE SOURCE: Prof. Dr. Z.-Y. Tang, Liver Cancer Institute, Zhongshan

Hospital, Fudan University, 136 Yi Xue Yuan Road, Shanghai

200032, China. zytang@srcap.stc.sh.cn

SOURCE: World Journal of Gastroenterology, (2002) 8/3 (385-392).

Refs: 119

ISSN: 1007-9327 CODEN: WJGAF2

COUNTRY: China

DOCUMENT TYPE: Journal; General Review FILE SEGMENT: 048 Gastroenterology

016 Cancer

029 Clinical Biochemistry

005 General Pathology and Pathological Anatomy

022 Human Genetics

LANGUAGE: English SUMMARY LANGUAGE: English

L21 ANSWER 13 OF 16 SCISEARCH COPYRIGHT 2003 THOMSON ISI on STN AB Protein-protein interactions play crucial roles in biological processes. Experimental methods have been developed to survey the proteome for interacting partners and some computational approaches have been developed to extend the impact of these experimental methods. Computational methods are routinely applied to newly discovered genes to infer protein function and plausible protein-protein interactions. Here, we develop and extend a quantitative method that identifies interacting proteins based upon the correlated behavior of the evolutionary histories of protein ligands and their receptors. We have studied six families of ligand-receptor pairs including: the syntaxin/Unc-18 family, the GPCR/G-alpha's, the TGF-beta/TGF-beta receptor system, the immunity/colicin domain collection from bacteria, the chemokine/chemokine receptors, and the VEGF/VEGF receptor family. For correlation scores above a defined threshold, we were able to find an average of 79% of all known binding partners. We then applied this method to find plausible binding partners for proteins with uncharacterized binding specificities in the syntaxin/Unc-18 protein and TGF-beta/TGF-beta receptor families. Analysis of the results shows that co-evolutionary analysis of interacting protein families can reduce the search space for identifying binding partners by not only finding binding partners for uncharacterized proteins but also recognizing potentially new binding partners for previously characterized proteins. We believe that correlated evolutionary histories provide a route to exploit the wealth of whole genome sequences and recent systematic proteomic results to extend the impact of these studies and focus experimental efforts to categorize physiologically or pathologically relevant protein-protein interactions. (C) 2002 Elsevier Science Ltd. All rights reserved.

ACCESSION NUMBER: 2002:986433 SCISEARCH

THE GENUINE ARTICLE: 620JP

TITLE: Co-evolutionary analysis reveals insights into

protein-protein interactions

AUTHOR: Goh C S; Cohen F E (Reprint)

CORPORATE SOURCE: Univ Calif San Francisco, Program Biol & Med Informat, San

Francisco, CA 94143 USA (Reprint); Univ Calif San Francisco, Dept Cellular & Mol Pharmacol, San Francisco, CA 94143 USA; Univ Calif San Francisco, Dept Biochem &

Biophys, San Francisco, CA 94143 USA

COUNTRY OF AUTHOR:

USA SOURCE:

JOURNAL OF MOLECULAR BIOLOGY, (15 NOV 2002) Vol. 324, No.

1, pp. 177-192.

Publisher: ACADEMIC PRESS LTD ELSEVIER SCIENCE LTD, 24-28

OVAL RD, LONDON NW1 7DX, ENGLAND.

ISSN: 0022-2836. Article; Journal

DOCUMENT TYPE: LANGUAGE:

English

REFERENCE COUNT:

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

L21 ANSWER 14 OF 16 WPIDS COPYRIGHT 2003 THOMSON DERWENT on STN

2001-589807 [66] ANWPIDS

WO 200162965 A UPAB: 20011113 AB

> NOVELTY - Differential expression screening to identify a genetic element involved in a cellular process (CP), involves comparing gene expressions in two cells (C1 and C2), where C2 has altered levels of a biological molecule implicated in CP, and identifying a genetic element whose expression differs, where gene expression in C1 or C2 is compared under different environmental conditions related to CP.

> DETAILED DESCRIPTION - Differential expression screening for identifying a genetic element involved in a cellular process, involves comparing gene expression in (C1) and gene expression in (C2), where C2 comprises altered levels, relative to physiological levels, of a biological molecule implicated in the cellular process, due to the introduction of a heterologous nucleic acid directing expression of a polypeptide, into C2 and identifying a genetic element whose expression differs, where gene expression in C1 or C2 is compared under at least two different environmental conditions relevant to the cellular process.

INDEPENDENT CLAIMS are also included for the following:

- (1) differential expression screening for identifying a gene product involved in a disease process, involving comparing gene expression in C1 and C2, and comparing gene expression in C1 and a third cell of interest (C3) that comprises altered levels, relative to physiological levels, of a candidate gene product, due to the introduction of a heterologous nucleic acid directing expression of the candidate gene product into C1, and selecting those candidate gene products which give rise to an alteration in the levels of expression of a second gene product in C3 relative to C1, where the second gene product also has altered levels of expression in C2 relative to C1; and
- (2) increasing the sensitivity of differential expression screening in which gene expression of C1 and C2 in response to two different levels of a signal are compared, by introducing a heterologous nucleic acid into C1 or C2 to increase the level of a biological molecule which modulates the response of the cell to the signal.

USE - The method is useful for differential expression screening of a genetic element involved in a cellular process (claimed). The method is also useful for identifying mutations and polymorphisms that affect the biological response to a particular cellular process. The method also allows the molecular dissection of biological pathways by altering a particular pathway under study.

ADVANTAGE - The method is an improved screening technique based on differential expression of genes, and allows the identification of other elements that are associated with genes that are implicated in a particular cellular process. By influencing the level of a particular biological molecule that is implicated in the pathway under study, through the introduction of the heterologous nucleic acid into one cell population, the method allows a pathway to be dissected into its precise molecular components.

Dwg.0/12

ACCESSION NUMBER:

2001-589807 [66] WPIDS

DOC. NO. CPI:

TITLE:

C2001-174860

Screening a genetic element involved in a cellular process, comprises comparing gene expressions in

a cell, and a second cell that has

altered levels of genes used in the process,

and detecting an element with varied expression.

DERWENT CLASS: B04 D16

INVENTOR(S):

KINGSMAN, A J

PATENT ASSIGNEE(S):

(OXFO-N) OXFORD BIOMEDICA UK LTD

COUNTRY COUNT:

PATENT INFORMATION:

PATENT NO KIND DATE WEEK PG ______

WO 2001062965 A2 20010830 (200166)* EN 103

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ

NL OA PT SD SE SL SZ TR TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM

DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC

LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE

SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

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CN 1425075 A 20030618 (200358)

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PATENT NO	KIND	APPLICATION	DATE
WO 2001062	965 A2	WO 2001-GB758	20010222
AU 2001033	937 A	AU 2001-33937	20010222
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		WO 2001-GB758	20010222
CN 1425075	Α	CN 2001-808359	20010222

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AU	200103393	37 A	Based	on	WO	2001	062965
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PRIORITY APPLN. INFO: GB 2000-18679 20000728; GB 2000-4197 20000222

L21 ANSWER 15 OF 16 SCISEARCH COPYRIGHT 2003 THOMSON ISI on STN The advances in cell culture engineering science have played AΒ a major role in the rapid expansion of mammalian cell-based products in the last decade. The host cell lines frequently used in manufacturing have converged to only a few species. For some, the genome has been completely sequenced. For others, the genomic sequence of closely related species is available, or will soon be available. The genomic and proteomic research tools enable one to globally survey the alterations at mRNA and protein levels to discover both primary and collateral gene expression changes and to unveil their regulation. Undoubtedly, a better understanding of these cellular processes

at the molecular level will lead to a better strategy for "designing"

producing cells and for process optimization. Herein

the genomic and proteomic tools are briefly reviewed and their

impact on cell culture Engineering is discussed.

ACCESSION NUMBER: 2001:5

2001:530691 SCISEARCH

THE GENUINE ARTICLE: 447FA

TITLE: Genomic and proteomic approaches in mammalian

cell culture technology
Korke R; Hu W S (Reprint)

AUTHOR: Korke R; Hu W S (Reprint)

CORPORATE SOURCE: Univ Minnesota, Dept Chem Engn & Mat Sci, 421 Washington

Ave SE, Minneapolis, MN 55455 USA (Reprint); Univ

Minnesota, Dept Chem Engn & Mat Sci, Minneapolis, MN 55455

USA USA

COUNTRY OF AUTHOR:

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JOURNAL OF THE CHINESE INSTITUTE OF CHEMICAL ENGINEERS,

(MAY 2001) Vol. 32, No. 3, pp. 213-218.

Publisher: CHINESE INST CHEMICAL ENGINEER, NATL TSING HUA UNIV, HSINCHU, DEPT CHEMICAL ENGINEERING, TAIPEI 300,

TAIWAN.

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L21 ANSWER 16 OF 16 SCISEARCH COPYRIGHT 2003 THOMSON ISI on STN

AB Hepatic stellate cell activation is a complex process

. Paradoxes and controversies include the origin(s) of hepatic stellate

cells, the regulation of membrane receptor signaling and transcription, and the fate of the cells once liver injury

resolves. Major themes have emerged, including the dominance of autocrine signaling and the identification of counterregulatory stimuli that oppose

key features of activated cells. Advances in analytical methods

including proteomics and gene array, coupled with powerful

bioinformatics, promise to revolutionize how we view cellular responses.

Our understanding of stellate cell activation is likely to

benefit from these advances, unearthing modes of regulating cellular

behavior that are not even conceivable on the basis of current paradigms.

ACCESSION NUMBER: 2000:546388 SCISEARCH

THE GENUINE ARTICLE: 333RN

TITLE: Fibrogenesis I. New insights into hepatic stellate

cell activation: the simple becomes complex

AUTHOR: Eng F J; Friedman S L (Reprint)

CORPORATE SOURCE: CUNY MT SINAI SCH MED, DEPT MED, DIV LIVER DIS, BOX 1123,

1425 MADISON AVE, RM 1170F, NEW YORK, NY 10029 (Reprint); CUNY MT SINAI SCH MED, DEPT MED, DIV LIVER DIS, NEW YORK,

NY 10029

COUNTRY OF AUTHOR:

USA

SOURCE:

AMERICAN JOURNAL OF PHYSIOLOGY-GASTROINTESTINAL AND LIVER

PHYSIOLOGY, (JUL 2000) Vol. 279, No. 1, pp. G7-G11. Publisher: AMER PHYSIOLOGICAL SOC, 9650 ROCKVILLE PIKE,

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